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(54) NOVEL SUGAR-CHAIN SYNTHETASE AND PROCESS FOR PRODUCING THE SAME

(57) Novel GalNAcα2,6-sialyltransferases P-B1 and P-B3; GalNAcα2,6-sialyltransferase genes encoding the above GalNAcα2,6-sialyltransferases P-B1 and P-B3; and an extracellularly releasable protein catalyzing GalNAcα2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAcα2,6-sialyltransferase P-B1 or P-B3 together with a signal peptide are provided. Also provided is a process for preparing a sialyltransferases which enables efficient recovery of a sialyltransferase expressed in a large quantity in microorganisms.

Description

Technical Field

The present invention relates to a sugar-chain synthetase and a DNA encoding the enzyme. More specifically, the present invention relates to an N-acetylgalactosamineo2,6-sialyltransferase (GalNAco2,6-sialyltransferase) and a DNA encoding the enzyme. The enzyme is useful as medicaments having inhibitory activities against tumor metastases and viral infection, and as agents for introducing a sialic acid moieties into drugs to increase their biological activity.

The present invention further relates to a process for producing the sugar-chain synthetase. More specifically, the present invention relates to a process for expressing sialyltransferases in microorganisms to obtain the sialyltransferases in large quantities.

Background Art

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Sialic acids play an important role in a variety of biological processes, like cell-cell communication, cell-substrate interaction, adhesion. It has been known that various kinds of distinguishable cell surface sialic acids exist which change in a regulated manner during development, differentiation, and oncogenic transformation.

Sialic acids occur at the terminal positions of the carbohydrate groups of glycoproteins and glycolipids, and they are enzymatically introduced from CMP-Sia to these positions in a post translational process. For example, three linkage patterns, Siao2,6Gal, Siao2,3Gal and Siao2,6GalNAc are commonly found in glycoproteins (Hakomori, S., Ann. Rev. Biochem., 50, pp.733-764, 1981), and two, Siao2,3Gal and Siao2,8Sia, occur frequently in gangliosides (Fishman, P., and Brady, R.O., Science, 194, pp.906-915, 1976).

The enzymes responsible for such enzymatic introduction of sialic acid (sialic acid transfer) as mentioned above are glycosyltransferases called sialyltransferases. It has been known that at least 12 different sialyltransferases are required to synthesize all known sialyloligosaccharide structures (Broquet, P. et al., Int. J. Biochem., 23, 385-389, 1991; and Weinstein, J. et al., J. Biol. Chem., 262, 17735-17743, 1987). Among these enzymes, five sialyltransferases have been purified so far, and it has been known that they exhibit strict specificity for acceptor substrate (Sadler, J. et al., J. Bio. Chem., 254, pp.4434-4443, 1979; Weinstein, J. et al., J. Biol. Chem., 257, pp.13835-13844, 1982; Rearick, J. et al., J. Biol. Chem., 254, pp.4444-4451, 1979; and Joqiasse, D.H. et al., J. Biol. Chem., 260, 4941-4951, 1985).

As for cDNAs encoding the aforementioned sialytransferases, cDNAs encoding Galβ1,4GlcNAcα2,6-sialytransferase (Galβ4GlcNAc-α6ST) have been cloned from various organs including liver (Weinstein, J. et al., J. Biol. Chem., 262, pp.17735-17743, 1987; Grundmann U. et al., Nucleic Acids Res. 18, 667, 1990; Bast, B. et al., J. Cell. Biol., 116, pp.423-435, 1992; and Hamamoto, T. et al., Bioorg. and Medic. Chem., 1, pp.141-145, 1993). Furthermore, cDNAs encoding Galβ1,3GalNAcα2,3-sialytransferase (Galβ3GalNAc-α3ST) (Gillespie, W. et al., J. Biol. Chem., 267, pp.21004-21010, 1992; Japanese Patent Unexamined Publication No. 5-504678/1993; and Lee, Y. et al., Eur. J. Biochem, 216, 377-385, 1993); Galβ1,3(4) GlcNAcα2,3-sialytransferase (Galβ3(4)GlcNAc-α3ST) (Wen, D. X et al., J. Biol. Chem., 267, 21011-21019,1992; and Kitagawa, H. et al., Biochem. Biophys. Res. Commun. 194, 375); and Galβ1,3GalNAc/Galβ1,4GlcNAcα2,3-sialytransferase (Sasaki, K. et al., J. Biol. Chem., 268, 22782-22787, 1993) have also been cloned.

With respect to GalNAco2,6-sialyltransferase, the isolation of this enzyme has been reported (Hakomori, S., Ann. Rev. Biochem., 50, 733-764, 1981). However, the enzyme has not been purified so as to be characterized as a single identifiable substance, and accordingly, the enzyme has not been practically used because of insufficient reaction specificity, stability, and quantitative availability. Furthermore, a cDNA sequence encoding GalNAco2,6-sialyltransferase (EC 2.4.99.3; GalNAc-α6ST) has not yet been cloned.

Each of the aforementioned sialyttransferases whose structures having been revealed has a hydrophobic segment located at the NH₂-terminal region, and is a type II transmembrane protein immobilized to cell membrane by the hydrophobic segment. From this reason, a problem arises that expressed enzymes are immobilized to cell membranes and are not capable of being extracellularly released, where expressions are carried out using vectors containing sialyl-transferase genes that are transfected into mammalian cells. Furthermore, another problem may arise, when the expression is performed using mammalian cells, that enzyme expressions may be reduced as endoplasmic enzyme concentrations exceed certain levels.

In order to solve the above problems, an extracellularly releasable fused protein may be prepared which comprises an active domain of a sialyltransferase and a signal peptide region. This method is characterized in that a sialyltransferase can be readily recovered from a cell cultivation mixture, because the method involves the step of extracellular release of the fused protein which retains sialyl transfer activity and function as a sialyltransferase. However, where the expression of a sialyltransferase is performed using a mammalian cell, a transfected cell may be unstable or trouble-some cultivation procedures are required. In addition, in order to obtain a large quantity of expressed sialyltransferase, a mass cell culture is essential for a long period of time, which may cause disadvantageous from viewpoints of cost and manufactural installations.

Processes are well known to those skilled in the art to obtain cloned cDNA sequence encoding an enzyme expressed in mammalian cells and prepare a recombinant vector containing a gene encoding the enzyme, per se, or in a soluble form, and to transform microorganisms with the vector. A desired enzyme can be produced, in a large quantity, by culturing the transformant obtained by the aforementioned method to allow the microorganism expresses the enzyme, per se, or in a soluble form that has the desired activity.

This process comprises, for example, a step of culturing a transformed microorganism and extracting an expressed enzyme by tysis of the microorganisms using lysozyme or the like. However, a large amount of insoluble or soluble proteins is expressed in the microorganisms in a short period of time, and such proteins may aggregate inside the microorganisms to form proteinic aggregates or precipitates. Accordingly, it is necessary to extract the protein from such aggregates or precipitates.

To extract the desired protein from the aforementioned aggregates or precipitates, generally employed methods are those using urea, guanidine hydrochloride and the like. In this approach, the expressed protein is generally subjected to denaturation using, for example, urea for solubilization (by an exposure of the hydrophobic region), and then to renaturation treatment. The renaturation may be achieved by removing the urea through dialysis. However, for the removal of urea, a problem is that optimal conditions including pH, salt concentration, and temperature must be chosen that are strictly specific to each of the enzymes, and this optimization of conditions is extremely time-consuming. If inappropriate conditions are applied, recovered enzyme may retain almost no activity, and therefore, the selection of the conditions for the renaturation is particularly important.

Accordingly, one object of the present invention is to provide purified GalNAco2,6-sialyltransferase. Another object of the present invention is to provide a DNA sequence encoding GalNAco2,6-sialyltransferase and an amino acid sequence of the enzyme by cloning a cDNA sequence that encodes GalNAco2,6-sialyltransferase. Further objects of the present invention are to provide an extracellularly releasable protein comprising an active domain of the GalNAco2,6-sialyltransferase and to provide a process for a mass expression of said protein in microorganisms. It is also an object of the invention to provide a process for extraction of an expressed sialyltransferase from aggregate thereof in microorganisms and a process of efficient renatuation of the extract.

SUMMARY OF THE INVENTION

The present inventors conducted various studies to achieve the foregoing objects, and as a result, they succeeded in cloning the cDNA encoding GalNAcα2,6-sialyltransferase from chick embryo. The present invention was achieved on the basis of these findings. The present invention thus provides GalNAcα2,6-sialyltransferase P-B1 characterized by the amino acid sequence disclosed as SEQ ID NO.1 in the sequence listings. The present invention also provides GalNAcα2,6-sialyltransferase genes encoding the aforementioned amino acid sequence of GalNAcα2,6-sialyltransferase P-B1, and as an embodiment thereof, a GalNAcα2,6-sialyltransferase gene characterized by the nucleotide sequence from nucleotide No.1 to 1698 disclosed as SEQ ID NO.1 in the sequence listings. Also provided are recombinant vectors comprising the above GalNAcα2,6-sialyltransferase gene and plasmid λCEB-3034 as an embodiment thereof, transformants which are transformed with the above recombinant vector, and the active domain of GalNAcα2,6-sialyltransferase characterized by the amino acids of No. 233 through 566 of the amino acid sequence disclosed as SEQ ID NO.1 in the sequence listings.

The GalNAc α 2,6-sialyltransferase P-B1 has activity of transferring sialic acid to the 6-position of N-acetylgalactosamine directly bound to a protein regardless of the presence or absence of a substituent on hydroxyl group at the 3position. The structure of NeuAc α 2,6GalNAc-protein is thus readily formed by the enzyme, which terminates further extension of the resulting sugar chain. Therefore, where a longer sugar chain is desired, a sugar chain synthetic scheme should be designed so that this enzyme can be employed after complete extension of a sugar chain. For this reason, a sialyltransferase is highly useful which fails to transfer sialic acid to an N-acetylgalactosamine that has unsubstituted 3-hydroxyl group and bonded to a protein via an α -glycoside linkage, but can transfer sialic acid to the 6-position of an N-acetylgalactosamine bound to a protein via an α -glycoside linkage, only when the hydroxyl group at 3-position is substituted with a galactose or a sugar chain having a galactose at its reduced terminus.

Therefore, the inventors of the present invention cloned a cDNA from chicken testes that encodes GalNAcα2,6-sialytransferase having the aforementioned features, and as a result, they achieved the present invention relating to the GalNAcα2,6-sialytransferase P-B3 characterized by the amino acid sequence disclosed as SEQ ID NO.3 in the sequence listings. The present invention thus provides GalNAcα2,6-sialytransferase genes encoding the above amino acid sequence of the GalNAcα2,6-sialytransferase P-B3, and as an embodiment thereof, the GalNAcα2,6-sialytransferase e gene having the nucleotide sequence of from nucleotide No.1 to 1212 as disclosed as the SEQ ID NO.3 in the sequence listings. The present invention also provides a recombinant vector comprising the above GalNAcα2,6-sialytransferase gene and plasmid λCEB3-T20 as an embodiment thereof, and a transformant being transformed with the above recombinant vector.

The inventor of the present invention further conducted studies to provide an extracellularly releasable protein comprising a portion, i.e. active domain, that is derived from the structure of the aforementioned GalNAco2,6-sialyltrans-

ferase and is responsible for its activity. As a result, they succeeded in identifying a partial polypeptide of the above-described GalNAco2,6-sialyltransferase as being the active domain, and achieved the present invention directed to an extracellularly releasable protein which comprises the polypeptide region together with a signal peptide and catalyzes GalNAco2,6-sialic acid transfer. As an embodiment thereof, protein SB-690 characterized by the amino acid sequence disclosed as SEQ ID NO.2 in the sequence listings. The present invention also provides genes encoding the above protein, and as an embodiment thereof, a gene having the nucleotide sequence characterized by from nucleotide No.1 to 1065 disclosed as SEQ ID NO.2 of the sequence listings, and a recombinant vector containing the aforementioned gene and plasmid pcDSB-690 as an embodiment thereof. Further provided are a transformant being transformed with the above recombinant vector and a process for preparing the aforementioned protein which comprises the steps of culturing the above transformant and recovering the above protein from the culture.

In addition, the inventors found that a Galβ1,4GalNAcc2,6-sialyltransferase with a highly restored activity can be prepared by expressing mouse Galβ1,4GalNAcc2,6-sialyltransferase in an insoluble form in Escherichia coli, followed by extracting the enzyme with urea and subjecting the enzyme to renaturation under optimal conditions, and thus achieved the present invention. In accordance with the present invention, there is provided a process for producing a sialyltransferase which comprises the steps of: (a) expressing a sialyltransferase in a microorganism; (b) extracting the sialyltransferase with about 5 to 9 M urea from proteinic aggregates or precipitates accumulated inside the microorganism and containing the enzyme; (c) diluting the extract obtained by the above step (b) with a renaturation composition to obtain a primary dilution containing about 1 to 4 M urea; (d) diluting the primary dilution obtained by the above step (c) with a renaturation composition to obtain a secondary dilution containing about 0.5 to 2 M urea; and (e) removing urea from the secondary dilution obtained by the above step (d) by dialysis to afford a renatured sialyltransferase.

BRIEF DESCRIPTION OF THE DRAWINGS

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Fig. 1 shows a restriction map of the cDNA done encoding GalNAcα2,6-sialyltransferase P-B1. In the figure, E represents EcoRI; RV: EcoRV; P: Pstl; and B: BgIII.

Fig. 2 shows the result of hydrophobicity analysis of the GalNAcα2,6-sialyltransferase P-B1 according to the present invention. In the figure, N-terminus of the protein is depicted at the left side and positive values indicate hydrophobic regions.

Fig. 3 shows the location of the active domain of the GalNAcα2,6-sialyltransferase P-B1 according to the present invention and the result of comparison with the structure of protein SB-690 which has GalNAcα2,6-sialyltransferase activity and can be extracellularly released. In the figure, protein SB-BGL is a protein not having GalNAcα2,6-sialyltransferase activity.

Fig. 4 shows the result of comparison between the primary sequences of GalNAco2,6-sialyltransferase P-B3 and GalNAco2,6-sialyltransferase P-B1 according to the present invention. In the figure, amino acids are represented by the one-letter abbreviations.

BEST MODE FOR CARRYING OUT THE INVENTION

As the most preferred embodiments of the present GalNAco2,6-sialyltransferases, GalNAco2,6-sialyltransferases P-B1 and P-B3 are provided. The explanations set out below will detail GalNAco2,6-sialyltransferases P-B1 and P-B3 as examples of the enzyme of the present invention. However, the GalNAco2,6-sialyltransferases of the present invention are not limited to the GalNAco2,6-sialyltransferase P-B1 or P-B3. GalNAco2,6-sialyltransferases comprising the active domain of the GalNAco2,6-sialyltransferase P-B1 and/or that of P-B3, both were first revealed by the present invention, or alternatively, those comprising one or more active domains of the GalNAco2,6-sialyltransferase in which the aforementioned acid sequence is partially changed or modified also fall within the scope of the present invention. A preferred example of such active domains as mentioned above is the active domain of the GalNAco2,6-sialyltransferase characterized by from amino acid No.233 to 566 of the amino acid sequence disclosed as SEQ ID NO.1 of the sequence listings.

The methods for isolation of the respective cDNAs encoding GalNAco2,6-sialytransferase P-B1 and GalNAco2,6-sialytransferase P-B3 will be detailed in Examples set out below. However, the methods for isolation of the cDNAs encoding GalNAco2,6-sialytransferase P-B1 and GalNAco2,6-sialytransferase P-B3 are not limited to those methods. One of ordinarily skilled artisan can readily isolate the desired cDNAs by referring to the methods described in the following examples, or alternatively, by appropriately modifying or altering those methods. In addition, the nucleotide sequences disclosed as SEQ ID Nos.1 through 3 in the sequence listings may be synthetically prepared and used to carry out the present invention.

The DNA sequence encoding GalNAco2,6-sialytransferase P-B1 as defined by SEQ ID No.1 in the sequence listings and the DNA sequence encoding GalNAco2,6-sialytransferase P-B3 as defined by SEQ ID No.3 are the preferred embodiments of the present invention. However, the DNA sequences encoding GalNAco2,6-sialytransferase P-B1 or GalNAco2,6-sialytransferase P-B3 of the present invention are not limited to those specified embodiments, and any

one of DNA sequences encoding the respective amino acid sequences of GalNAco2,6-sialyltransferase P-B1 and GalNAco2,6-sialyltransferase P-B3 revealed by the present invention fall within the scope of the present invention. For example, the DNA sequence encoding the active domain of GalNAco2,6-sialyltransferase characterized by the amino acids of from No. 233 to 566 of the amino acid sequence as defined by SEQ ID No.1 in the sequence listings is a preferred embodiment of the present invention. In addition, the DNA characterized by the nucleotides sequence of from nucleotide No. 699 to 1698 of the SEQ ID No.1 shown in the sequence listings is a particularly preferred embodiment of the present invention.

The GalNAca2,6-sialyltransferases of the present invention, including P-B1 and P-B3 for example, may occasionally be retained inside the cells after expression and not released extracellularly. Furthermore, when endoplasmic concentrations of the enzymes exceed certain levels, expressed amounts of the enzymes may possibly be reduced. In order to efficiently utilize the aforementioned GalNAca2,6-sialic acid transfer activities of GalNAca2,6-sialyltransferase P-B1 and P-B3, proteins in soluble forms may be prepared in which the activities of these enzymes are retained and can be released extracellularly from cells upon their expressions. Examples of such proteins include, for example, extracellularly releasable proteins which comprise a polypeptide, as being an active domain of the above-described GalNAca2,6-sialyltransferase P-B1 or P-B3 and is responsible for the GalNAca2,6-sialyltransferase activity, and catalyze the GalNAca2,6-sialic acid transfer.

The sialyltransferases so far cloned have domain structures similar to those of other glycosyltransferases: a short NH₂-terminal cytoplasmic tail; a hydrophobic signal-anchor domain; a proteolytically sensitive stem region; and a large COOH-terminal active domain (Paulson, J.C. and Colley, K.J., J. Biol. Chem., 264, 17615-17618, 1989). To determine the location of the transmembrane domain of the GalNAco2,6-sialyltransferase P-B1 of the present invention, hydropathy plot may be used which can be prepared according to the method of Kyte and Doolittle (Kyte, J. and Doolittle, R.F., J. Mol. Biol., 157, 105-132, 1982). To evaluate a putative active domain, recombinant plasmids introduced with various fragments may be produced and utilized. Exemplary methods will be detailed in the Examples set out below. However, the methods for determination of the location of the transmembrane domain or evaluation of a putative active domain are not limited to the disclosed methods.

For the preparation of the extracellularly releasable protein comprising a polypeptide portion, as being an active domain of the above-described GalNAca2,6-sialyltransferase P-B1 or P-B3, together with a signal peptide, an immunoglobulin signal peptide sequence, for example, may be used as the signal peptide, and a sequence corresponding to the active domain of GalNAca2,6-sialyltransferase P-B1 or P-B3 may be fused in-frame to the signal peptide. For example, the method of Jobling et al. (Jobling, S.A. and Gehrke, L., Nature (Lond.), 325, 622-625, 1987) may be applied as such methods, whose specified procedure will be detailed in the Examples of the present specification with respect to GalNAca2,6-sialyltransferase P-B1. However, types of the signal peptide and methods for ligation of the signal peptide and the active domain are not limited to the aforementioned methods, and a person skilled in the art can suitably choose the polypeptide portion as being an active domain of GalNAca2,6-sialyltransferase, preferably GalNAca2,6-sialyltransferase P-B1 or P-B3, and produce the extracellularly releasable protein by ligating the polypeptide portion to any available signal peptide according to an appropriate method. The most preferred example of these proteins is protein SB-690 of the present invention.

According to another embodiment of the present invention, there is provided a process for producing a sialyltransferase which comprises the steps of: (a) expressing a sialyltransferase in a microorganism; (b) extracting the sialyltransferase with about 5 to 9 M urea from proteinic aggregate or precipitate containing the enzyme and being accumulated inside the microorganism; (c) diluting the extract obtained by the above step (b) with a renaturation composition to obtain a primary dilution containing about 1 to 4 M urea; (d) diluting the primary dilution obtained by the above step (c) with a renaturation composition to obtain a secondary dilution containing about 0.5 to 2 M urea; and (e) removing urea from the secondary dilution obtained by the above step (d) by dialysis to afford a renatured sialyltransferase. As described above, sialyltransferases share the common domainal structure, and therefore, the preparation process of the present invention may be applicable to any type of sialyltransferase. For example, GalNAco2,6-sialyltransferase of Galβ1,4GalNAco2,6-sialyltransferase of the present invention can be suitably prepared by the process of the present invention.

According to an embodiment of the process of the present invention, 8 M urea is used in the step (b); a primary dilution containing about 2 to 3 M urea is obtained in the step (c); a secondary dilution containing about 1 to 2 M urea is obtained in the step (d); and the secondary dilution is dialyzed in the presence of divalent cations in the step (e). According to another embodiment of the present method, 8 M urea is used in the step (b); a primary dilution containing about 2 to 3 M urea is obtained by being left stand for 12 hours or more at 4 °C after primary dilution in the step (c); a secondary dilution containing about 1 to 2 M urea is obtained by being left stand for 48 hours or more after secondary dilution in the step (d); and the secondary dilution is dialyzed in the presence of divalent cations in the step (e). In addition, it is also a preferred method in which the renaturation composition used in the step (c) contains 1 to 2 M urea, 20 mM MOPS-NaOH, 0.5M NaCl, 20 mM lactose, 0.5 mM EDTA (pH 7.0) and the renaturation composition used in the step (d) contains 20 mM MOPS-NaOH, 0.5M NaCl, 20 mM lactose, 0.5 mM EDTA (pH 7.0).

The first step of the process for the preparation of sialyttransferase according to the present invention is the expression of a sialyttransferase in microorganisms. To this end, previously doned genes of sialyttransferases can be used. As cDNAs encoding sialyttransferases, the cDNA encoding Galβ1,4GlcNAcα2,6-sialyttransferase (Galβ4GlcNAc-α6ST, see, Weinstein et al., Grundmann et al., Bast et al. and Hamamoto et al., supra), the cDNA encoding Galβ1,3(4)GlcNAcα2,3-sialyttransferase (Galβ3(4)GlcNAc-α3ST, see, Wen et al. and Kitagawa et al., supra), the cDNA encoding Galβ1,3GalNAc/Galβ1,4GlcNAcα2,3-sialyttransferase (see, Sasaki et al., supra), the cDNA encoding Galβ1,3GalNAcα2,3-sialyttransferase (Galβ3GalNAc-α3ST, see, Gillespie et al. and Japanese Patent Unexamined Publication No. 5-504678/1993; and Lee et al., supra), for example, may be used, as well as cDNAs encoding the GalNAcα2,6-sialyttransferases of the present invention. Sialytransferase genes contained in these nucleotide sequences, per se, may be used for the expression of the naturally-derived enzymes.

According to the present invention, in addition to the naturally-derived sialyltransferases mentioned above, non-natural sialyltransferases in which the polypeptide sequences of the naturally-derived sialyltransferases are partly deleted or modified may be expressed in microorganisms. For example, since sialyltransferases have a hydrophobic segment (transmembrane domain) in the NH₂-terminal region, and sialyltransferases in soluble forms wherein the hydrophobic segment is deleted are preferably expressed in the microorganisms. In addition, deletion of both of the hydrophobic segment and the cytosol segment is also preferred.

In order to produce recombinant vectors for the expression of sialyltransferases, the entire sequences or partial regions of the genes of naturally derived sialyltransferases may be selectively amplified by, for example, PCR method. For example, a sialyltransferase gene (a PCR fragment) may be readily prepared which has an initiation codon and a cloning site and lacks the cytosol domain and transmembrane domain. This type of sialyltransferase genes are suitably used for the introductions into vectors for microbial expressions due to the presence of the initiation codon and the cloning site. In addition, said genes are preferred since they encode non-natural sialyltransferases, in which a part of the polypeptide sequence of the naturally-derived sialyltransferase is deleted, and express non-natural soluble sialyltransferase in microorganisms.

According to the process of the present invention, microorganisms such as Escherichia coli may be used for the expression of sialyltransferase. A microbial expression vector suitably used for transformation of such microorganisms may be suitably selected by an ordinarily skilled artisan. For example, where E. coli JM109(DE3) or the like is used as the microorganism, microbial expression vectors such as pET3b (Studier, F.W. et al., Method. Enzymol., 185, pp.60-89, 1990) may be used. Methods for introducing the above described sialyltransferase genes into microbial expression vectors and methods for transforming microorganisms with recombinant vectors are both well known to those skilled in the art.

The transformants can be cultured according to methods for culturing transformed microorganisms well known to those skilled in the art. For efficient expression of a desired sialytransferase in microbial cells, replication of the recombinant protein can be initiated by, for example, the induction of T7-RNA polymerase during the logarithmic growth phase of the transformants. A large amount of naturally-derived or non-natural sialytransferase is expressed inside the transformants thus obtained, which generally forms proteinic aggregate or precipitate.

The second step of the process of the present invention is the extraction step of a sialyltransferase with 5 to 10 M urea from the proteinic aggregate or precipitate which is accumulated inside the cells and contains the sialyltransferase. In order to expose the proteinic aggregate or precipitate to outside of the microorganisms for its separation, the cultured transformants can be treated with, for example, lysozyme or Triton X-100 and then insoluble fractions may be collected by centrifugation. After then, the precipitates are suspended in a buffer (for example, 10 mM Tris-HCl, pH 7.4) at a protein concentration of about 1 to 10 mg/ml and are subjected to extraction with urea.

For example, solid urea is added to the suspension so as to be 5 to 10 M, preferably 8 M of final concentration, and the precipitates are subjected to extraction for 15 minutes to 2 hours, preferably 30 minutes at 4 to 25 °C, preferably at 10°C. While not bound by any specific theory, the hydrophobic portion of a sialyltransferase contained in the extract is exposed by the action of urea, and as a result, a solubilized sialyltransferase is extracted from the proteinic aggregates or precipitates.

Then, an extract solution containing a denatured sialyltransferase can be obtained by removing the precipitates by, for example, centrifugation of the extract at 12,000 x g for 15 minutes. This extract normally contains about 0.5 mg/ml of proteins. For example, when 5.7 M urea is used for the extraction, about 80% of proteins can be recovered. Furthermore, upon the extraction, NaCl and Tris-HCl (pH 7.4) are preferably added so that their final concentrations of 0.3 M and 20 mM, respectively, are achieved. Exemplary procedure of the extraction will be explained in detail in the Examples set out below.

The sialytransferase contained in the extract exposes hydrophobic portions and its higher-order structure is damaged. According to the process of the present invention, renaturation of the sialytransferase contained in the extract is performed as the third step. The term renaturation herein used means restoration of the higher-order structure of the protein that is lost during the extraction step and the entire or partial recovery of the enzymatic activity. This step is characterized in that the extract is diluted stepwise with a renaturation composition so that the urea concentration can be gradually lowered to efficiently achieve the renaturation of the sialytransferase.

The renaturation process comprises the steps of, for example, diluting the extract with a renaturation composition to obtain a primary dilution containing about 1 to 4 M urea; diluting the primary dilution with a renaturation composition to obtain a secondary dilution containing about 0.5 to 2 M urea; and removing the urea from the secondary dilution by dialysis to afford a renatured sialytransferase.

A preferred embodiment of the process comprises the steps of, for example, diluting the extract with a renaturation composition to obtain a primary dilution containing about 2 to 3 M urea; diluting the primary dilution with a renaturation composition to obtain a secondary dilution containing about 1 to 2 M urea; and removing the urea from the secondary dilution by dialysis in the presence of one or more divalent cations to afford a renatured sialyltransferase. A further preferred embodiment is a process comprises the steps of, for example, diluting the extract with a renaturation composition and the result is allowed to stand for 12 hours or more at 4°C to obtain a primary dilution containing about 2 to 3 M urea; diluting the primary dilution with a renaturation composition and the result is allowed to stand for 48 hours or more to obtain a secondary dilution containing about 1 to 2 M urea; and removing the urea from the secondary dilution by dialysis in the presence of one or more divalent cations to afford a renatured sialyltransferase.

As the renaturation composition, for example, 2 M urea, 20 mM MOPS-NaOH (MOPS: 3-morpholinopropanesulfonic acid) (pH 7.0), 0.5M NaCl, 10 mM lactose, 0.5 mM EDTA; and 2 M urea, 20 mM Tris-HCl, 0.3M NaCl, 20 mM lactose, 0.5 mM EDTA (pH 7.4) may be used. In addition, a modified composition may be used in which the components of the latter composition may be changed to, for instance, 20 mM Tris-HCl (pH 8.0); 20 mM MOPS-NaOH (pH 7.0); 20 mM MES-NaOH (pH 6.0) (MES: 3-morpholinoethanesulfonic acid); 0.5 M NaCl; 0.1 M NaCl; or 1M urea. Furthermore, compositions not containing urea or lactose may also be used. Among these, 2 M urea, 20 mM MOPS-NaOH, 0.5 M NaCl, 20 mM lactose, 0.5 mM EDTA (pH 7.0) is preferably used. When the concentration of NaCl is below 0.1 M, or pH exceeds 9, renaturation efficiency is undesirably reduced. Generally, a salt concentration of 0.3 to 0.5 M and pH of 6 to 8 are preferred after the addition of the renaturation composition.

The first dilution comprises the step of preparing a primary dilution using the aforementioned renaturation composition so that a final protein concentration of the extract is 0.01 to 0.05 mg/ml, preferably about 0.02 mg/ml. For example, the extract may be diluted 10 to 40-fold, preferably about 20-fold, and a urea concentration may be 1 to 4 M, preferably not higher than about 3 M and not lower than about 2 M. Dilution treatment is generally and preferably performed at 4°C. This primary dilution mixture is left stand for 12 hours or more at 4°C, most preferably for about 12 hours, to initiate gradual renaturation.

The secondary dilution is carried out by diluting the primary dilution with an equal volume of renaturation composition, preferably not containing urea, to achieve approximately the half urea concentration. Through this dilution, urea concentration of the secondary dilution should be lowered to about 0.5 to 2 M, preferably not higher than about 2 M and not lower than 1 M (e.g., 1 to 2 M), and most preferably at about 1.2 M. The secondary dilution is allowed to stand for 40 hours to 2 weeks, preferably 48 to 72 hours, most preferably about 48 hours at 4°C, to proceed gradual renaturation.

After then, to achieve perfect renaturation, the above-obtained secondary dilution is dialyzed against, for example, a renaturation composition free from urea to completely remove remaining urea. The dialysis may be carried out at 4 °C for about 48 hours. Dialysis solution may be, for example, any one of buffer solutions in which the sialystransferase can be stored stably, as well as the renaturation composition.

In addition, by carrying out the primary and secondary dilution and the final dialysis in the presence of one or more divalent cations, renaturation efficiency can be further improved. Examples of the divalent cations include, for example, magnesium ions and manganese ions. These ions may be used at a concentration of 1 to 10 mM, preferably about 5 mM. It is particularly preferred that the dialysis is performed in the presence of one or more divalent cations. When a reducing agent such as dithiothreitol and mercaptoethanol is added before complete removal of urea in the final dialysis step, the enzymatic activity may occasionally be lost. However, after the urea is completely removed, the enzyme restores resistance to the reducing agent to exhibit the sialyltransferase activity.

The present invention will be further explained more specifically by referring to the following examples. However, the scope of the present invention is not limited to these examples.

Examples

(A) Preparation of GalNAco2.6-sialytransferase P-B1

In order to obtain a cDNA clone of GalNAca2,6-sialyltransferase P-B1, PCR with two degenerate oligonucleotides (ST-107 and ST-205) was performed using chick embryo cDNA as a template. A fragment of the desired size of approximately 150 bp was obtained. Among the PCR recombinants, one clone, designated as CEB1, was found to have a unique amino acid sequence distinct from the known sialylmotifs of Galβ4GlcNAc-α6STRL (residues 180-225), Galβ3(4)GlcNAc-α3STRL (residues 158-203), and Galβ3GalNAc-α3STPS (residues 144-189). The homologies of the sialylmotif of CEB1 with those of Galβ4GlcNAc-α6STRL, Galβ3(4)GlcNAc-α3STRL and Galβ3GalNAc-α3STPS were 56%, 58% and 60%, respectively.

Screening of a 6-day-old chick embryo cDNA library with the cDNA insert from the CEB1 was carried out, and as a result, several cDNA clones were identified. Among them, clone λ CEB-3043 contained a 2.7 kb insert (Fig. 1). To obtain other overlapping clones, a random-primerd cDNA library was again screened by hybridization with the 0.8 kb EcoRl-Bglll fragment of the 5'-end of the λ CEB-3043. Fifteen clones were isolated from the cDNA library. Among them, one clone, λ CEBHAD contained a 220 bp insert overlapping with the 5'-end of clone λ CEB-3043 for 160 bp.

The combined DNA from these two cDNAs contained a 1.7 kb of open reading frame that ends at a TGA terminal codon at nucleotide 1699. A poly adenylation signal (AATAAA) at 23 nucleotides upstream from the poly(A) sequence exists at the 3'-end. Translation of this open reading frame affords GalNAcα2,6-sialyltransferase P-B1 of the present invention (occasionally referred to as P-B1 in the the examples) of 566 amino acids with a molecular mass of 64,781, which starts with a methionine codon at nucleotide 1 with a conventional initiation sequence (Kozak, M., Nature (Lond.), 308, 241-246, 1984). The cDNA including a gene encoding the GalNAcα2,6-sialyltransferase of the present invention, the nucleotide sequence of λCEB-3043 as being the gene encoding the GalNAcα2,6-sialyltransferase of the present invention, and the amino acid sequence of the GalNAcα2,6-sialyltransferase P-B1 of the present invention are shown in the SEQ ID No.1 of the sequence listings.

Polymerase chain reaction (PCR)

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PCR was performed using degenerate primers [5' primer ST107: TGGGCCTTGGII(A/C)AGGTGTGCTGTTG, and 3' primer ST205: AGGCGAATGGTAGTTTTTG(A/T)GCCCACATC] deduced from conserved regions in Galβ4GlcNAc-α6STRL (Weinstein, J. et al., J. Biol. Chem., 262, 17735-17743, 1987), Galβ4GlcNAc-α6STHP (Grundmann, U. et al., Nucleic Acids Res., 18, 667, 1990), and Galβ3GalNAc-α3STPS (Gillespie, W. et al., J. Biol. Chem., 267, 21004-21010, 1992). To obtain cDNA, poly(A)-rich RNA (2 μg) from 3 day-old chick embryos was incubated with an oligo-dT primer (Pharmacia), 1 mM each of dATP, dCTP, dGTP and dTTP, and 2 U/μl of RNase inhibitor (Promega) in 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.5 mM MgCl₂ and 0.001% gelatin in 50 μl for 10 min at 0 °C, and then for further incubation was carried out for 60 min at 42°C after the addition of 100 μU Moloney murine leukemia virus reverse transcriptase (BRL).

After heating the reaction mixture at 94°C for 3 min, cDNA prepared from 0.2 µg of poly(A)-rich RNA was used for the PCR experiment in a mixture comprising 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.25 mM MgCl₂ 0.001% gelatin, 200 µM of each dATP, dCTP, dGTP and dTTP, 2U of Taq DNA polymerase (Promega), and 40 pmoles of each PCR primer in 50 µl. PCR amplification, 35 cycles, was carried out, each cycle consisting of denaturation at 96°C for 45 sec, annéaling at 50°C for 60 sec, and extension at 72°C for 60 sec. The PCR products were developed on a 3% agarose gel. The DNA fragment corresponding to 150 bp was eluted from the gel (Qiaex kit; Qiagen), blunt-ended and kinated, and then subcloned into the Smal site of pUC119, and finally sequenced.

Construction of a cDNA library

Total RNA was prepared from chick embryos (6-day-old) by the guanidinium thiocyanate method, followed by centrifugation in a 5.7 M CsCl solution (Sambrook, J., Molecular Cloning: a Laboratory Manual, 2nd ed. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y. 1989). Poly(A)-rich RNA was purified with oligotex-dT30 (Takara), and then employed for the construction of a cDNA library using λ ZAPII (Stratagene) and cDNA synthesis (Pharmacia) kits with an oligo-dT primer and random primers.

Screening of the cDNA library

The amplified cDNA library (1 x 10^6 plaques) was screened with the chick embryo PCR fragments. The plaque-transferred filters were hybridized with 32 P-radiolabeled DNA probes for 12 h at 65°C in 5 x SSC, 0.02% SDS, 5 x Denhardt's solution and 10 µg/ml denatured salmon sperm DNA, and then washed twice at 65°C for 20 min in 2 x SSC, 0.1% SDS. To obtain plasmids from the isolated phage clones, phagemid rescue was performed according to the manual of the manufacturer of the λ ZAPII cloning kit (Stratagene). cDNA inserts were excised directly as Bluescript plasmids. Plasmids were produced by the standard molecular cloning method according to Sambrook et al. (Sambrook, J. et al., Molecular Cloning: a Laboratory Manual).

DNA sequence analysis

The DNA sequences of the inserts were determined by the dideoxy-chain termination method (Sanger, F. et al., Proc. Natl. Acad. Sci. USA, 74, 5463-5467, 1977) using single-strand DNA as a template for T7-DNA polymerase. The sequencing reaction and electrophoresis were carried out using the AutoRead DNA sequencing kit and a DNA sequencer (Pharmacia). Single strand DNA was prepared from Escherichia coli XL-Blue (Stratagene) after superinfection with helper phage R408 (Stratagene). The sequence data were analyzed with a computer using PC/Gene (Teijin System Technology).

Northern and Southern blot analyses

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To confirm the existence of the gene, Southern blot analysis of chick genomic DNA was performed. Hybridization with the EcoRI cDNA insert of λ CEB-201 gave a single band for the DNA digested with EcoRI and BamHI, and two bands for the DNA digested with HindIII and SacI. This simple hybridization pattern indicates that the cloned cDNA is a single copy gene.

The transcription pattern during embryonic development was examined by Northern blot hybridization. Analysis of RNA from 6, 8 and 10 day-old chick embryos revealed two RNA species of 3.0 and 2.2 kb. The 3.0 kb transcript was abundant and constantly expressed during all embryonal stages. A low level of the 2.2 kb transcript was detected in 6 day-old embryos and its expression was decreased in 8 and 10 day-old embryos. The gene expression was analyzed using 10 µg poly(A)-rich RNA obtained from various chicken tissues: brain, heart, liver, lung, kidney, and testis. Very low levels of the 3.0 and the 4.0 kb transcripts was detected in testes, while almost no signals were detected in other tissues. The following description details each of the experiments.

For Northern blots, $5\,\mu g$ of denatured poly(A)-rich RNAs from chick embryo was size-fractionated on formaldehyde-agarose gels and then blotted onto Hybond N+ nylon membranes (Amersham). For Southern blots, $7.5\,\mu g$ of genomic DNA prepared from chick embryos was digested with restriction enzymes EcoRI, BamHI, HindIII and SacI, and then size-fractionated on 0.6% agarose gels. After electrophoresis, the gels were denatured (30 min) in $0.5\,N$ NaOH and $1.5\,M$ NaCI and neutralized (30 min) in $0.5\,M$ Tris-HCI (pH 7.5) and $1.5\,M$ NaCI, and then the DNA was transferred onto Hybond N+ nylon membranes. Both Northern and Southern filters were prehybridized in 50% formamide, $5\times SSC$, $5\times Denhardt's$, 0.5% SDS, and $10\,\mu g/mI$ denatured salmon sperm DNA at $37\,^{\circ}C$ for $1\,h$, and then hybridized with a ^{32}P -labelled DNA probe for $12\,h$ under the same conditions as for prehybridization. The probe applied was a $0.6\,kD$ EcoRI cDNA insert of λ CEB-201, which was labeled with a Multiprime Labeling System (Amersham). The filters were washed twice for $10\,m$ min at $65\,^{\circ}C$ in $2\times SSC$ and 0.1% SDS, followed by washing twice with $0.2\times SSC$ and 0.1% SDS at $65\,^{\circ}C$ for $30\,m$, and then exposed to Kodak XAR film for about one day at $-70\,^{\circ}C$.

The amino acid sequence of the sialytransferase P-B1 of the invention, which was revealed as described above, shows the following characteristic features that are not observed in sialytransferases so far known.

- (i) All of the sialyltransferases previously cloned are critical Type II membrane proteins. They have a domain structure similar to that of other glycosyl-transferases: a short NH₂-terminal cytoplasmic tail; a hydrophobic signal-anchor domain; a proteolytically sensitive stem region; and a large COOH-terminal active domain. On the other hand, the sialyltransferase P-B1 of the invention has a large stem region (or intermediate region).
- (ii) The sialyltransferase P-B1 of the invention has a PEST region (residues 233-258). It has been known that the amino acid sequences of proteins with intracellular half-lives of less than 2 hours contain one or more regions that are rich in proline, glutamic acid, serine, and threonine residues (referred to as PEST: Rogers, S. et al., Science, 234, 364-368, 1986). These PEST regions are generally flanked by clusters containing several positively charged amino acids. Other sialyltransferases previously known do not have this region.
- (iii) Two stretches of eight amino acids (SSSXVSTC) were found at residues 247-254 and 330-337. A search of the Genebank database for other proteins revealed no sequence similarity to this sequence.

Sialytransferases so far known exhibit remarkable tissue-specific expression, which seems to be correlated with the existence of cell type-specific carbohydrate structures (Paulson, J.C. and Colley, K.J., J. Biol. Chem., 264, pp.17615-17618, 1989). The results of Northern blotting indicates that the pattern of expression of sialyltransferase P-B1 apparently changes. The transcriptions of three different sizes of mRNAs (4.0, 3.0 and 2.2 kb) from the sialyltransferase P-B1 gene suggests that they are generated through alternative splicing and alternative promoter utilization mechanisms as observed for Galβ1,4GicNAco2,6-sialyltransferase (Galβ4GicNAc-α6STRL) and Galβ1,3(4)GicNAco2,3-sialyltransferases (Galβ3(4)GicNAc-α3STRL, Weinstein, J. et al., J. Biol. Chem., 262, 17735-17743, 1987; and Wen, D.X. et al., J. Biol. Chem., 267, 21011-21019, 1992). This hypothesis is supported by the results of Southern hybridization, which showed the existence of a single copy gene for sialyltransferase P-B1.

(B) Preparation of the soluble form protein SB-690

In order to utilize the GalNAco2,6-sialyltransfer activity of the GalNAco2,6-sialyltransferase P-B1 of the present invention, protein SB-690 in a soluble form was prepared which retains the activity of the present enzyme and is released from the cells upon expression.

The sialyttransferases so far cloned have a domain structure similar to that of other glycosyl-transferases: a short NH₂-terminal cytoplasmic tail; a hydrophobic signal-anchor domain; a proteolytically sensitive stem region; and a large COOH-terminal active domain. To determine the location of any transmembrane domain of GalNAco2,6-sialytransferase of the present invention, a hydropathy plot (Fig.2) was prepared from the translated sequence according to the method of Kyte and Doolittle (Kyte, J. and Doolittle, R.F., J. Mol. Biol., 157, 105-132, 1982). As as result, it is suggested

that a critical hydrophobic transmembrane domain of GalNAco2,6-sialyltransferase P-B1 of the present invention consists of 21 amino acid residues from the amino adid No.17 to 37.

As described above, the hydrophobic signal anchor domain of GalNAco2,6-sialytransferase is lacated from amino acid resudie No.17 to 37. Residues from 233 to 269 apparently contain certain essential residues for the enzymatic activity, because the media from cells transfected with pcDSB-BGL had no significant activity, while the protein (33 KDa) was synthesized in an in vitro translation/transcription system with pSB-BGL as a template. The active domain was thus deduced to be around 233-566 (Fig.3), which is a comparative size to that of other cloned sialytransferases. In order to produce the soluble protein containing the active domain described above, the sequence relating to the putative active domain of P-B1 was in-frame fused to the sequence of immunoglobulin signal peptide (Jobling, S.A. and Gehrke, L., Nature (Lond.), 325, 622-625, 1987). Details of the experiments are shown below.

A vector plasmid pUGS was constructed by replacing the Pstl-Xhol fragment of the p Bluescript SK(+) plasmid with a 117 bp of a synthetic DNA fragment. This fragment contains 43 bp of the 5'-untranslated leader sequence of Alfalfa Mosaic Virus (Jobling, S.A. and Gehrke, L., Nature (Lond.) 325, 622-625, 1987) with a synthetic Pstl site at the 5'-end, followed by the mouse immunoglobulin M heavy chain signal peptide sequence (57 bp) (Boersch-Supan, M.E. et al., J. Exp. Med. 161, 1272-1292, 1985) with a 17 bp of a synthetic EcoRI, BgIII and Xhol doning site at the 3'-end. The nucleotide

5'-CTGCAGGGTTTTTATTTTTAATTTTCTTTCAAATA

CTTCCACCATGAAATTCAGCTGGGTCATGTTCTTCCTGATGGCAGTGGTTACAGGGGTCAATTCAGAA

TTCCAGATCTCGAG-3'.

s sequence of this fragment is

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λCEB-3043 encoding GalNAco2,6-sialytransferase of the present invention was partially digested with EcoRV, and 1.8 kb fragment was subcloned into EcoRV site of pBluescript SK(+) to generate pCEB-1800. This clone lacks 0.8 kb of 3'-untranslated region of λCEB-3043. An active domain of GalNAco2,6-sialytransferase P-B1 was generated by PCR using the 5'-primer, 5'-AGGGCTGCTGAATTCACTGAGCCACAG-3' (nucleotides 679-708), with a synthetic EcoRI site at the middle of the primer and a 3' universal M13 sequencing primer and pCEB-1800 as a template. The PCR product was digested with EcoRI and XhoI, and then ligated into the EcoRI/XhoI site of pUGS to yield the plasmid pSB-690. In this plasmid, a sequence obtained by in-frame fusion of the 3'-end of the immunoglobulin signal sequence to the putative active domain of GalNAco2,6-sialytransferase P-B1 was contained. The fusion fragment was excised from pSB-690 with PstI and XhoI, and then inserted into the PstI/XhoI site of expression vector pcDSRα to yield pcDSB-690.

As a control, protein SB-BGL which lacks the active domain of GalNAco2,6-sialytransferase was produced as described below. pCEB-1800 and pUGS were digested with Bglll, and the protruding ends were filled by using the Klenow fragment of DNA polymerase. After heat denaturation of the Klenow fragment of DNA polymerase (at 94 °C for 20 min), these plasmids were digested with Xhol. The 1.0 kb fragment from pCEB-1800 was gel purified and subcloned into the blunt-ended Bglll/Xhol site of pUGS to yield pSB-BGL. The Pstl/Xhol fragment from pSB-BGL was subcloned into the Pstl/Xhol site of pcDSRa to generate pcDSB-BGL.

Expression of the above described protein was performed as follows. COS-7 cells were transiently transfected with 5 μg of plasmid DNA using the DEAE-dextran method (McCutchan, J.H. and Pagano, J.S. J. Natt. Cancer Inst. 41, pp.351-357,1968). The media were harvested after 48 h transfection and then concentrated 10 times on Centricon 30 filters (Amicon) for the enzyme assay. For metabolic labeling, COS cells (60-mm culture dish) were washed with Metree medium (Dulbecco's modified Eagle's medium and 2% fetal calf serum) (GIBCO) and then incubated for 1 h with the same media. The cells were pulse-labeled with 10 MBq/dish of Express ³⁵S protein labeling mix (Du Pont-New England Nuclear) in 1.5 ml of Met-free media for 2 h. These cells were then washed with Met-free media and chased for 5 h in media without Express-label. The media containing secreted proteins were harvested, concentrated 10 times, and then subjected to SDS-PAGE, followed by fluorography.

The enzyme activity of the expressed protein was measured as follows. The assays using oligosaccharides and glycoproteins as acceptors were performed in the presence of 50 mM sodium cacodylate buffer (pH 6.0), 50 µM CMP-[14C-]NeuAc (0.9 Bq/pmol), 1 mg/ml of bovine serum albumin, 2 mg/ml of acceptor substrate and 1 µl of concentrated COS cell medium, in a final volume of 10 µl, and were incubated at 30 °C for 2 h. At the end of the incubation period, 1 µl of the assay mixture was applied to a Silica gel 60HPTLC plate (Merck, Germany). The plate was developed with ethanol:pyridine:n-butanol:water:acetate (100:10:10:30:3), and the radioactivity was visualized and quantified with a BAS2000 radio image analyzer (Fuji Photo Film, Japan). The radioactivity remaining at the origin was taken as sialylated glycoprotein.

Identification of the sialylated products was carried out as follows. Asialo-BSM were resialylated with CMP[14C]NeuAc in pcDSB-690 COS cell medium and β-elimination oligosaccharides were prepared. β-elimination was car-

ried out according to Carlson's method (Carlson, D.M., J. Biol. Chem., 243, 616-626, 1968). Asialo-BSM (100 μ g each) was sialylated with CMP-[14 C]NeuAc in pcDSB-690 COS cell medium under the same conditions as above, except that the incubation period was 12 h. The reaction was terminated by adding 500 μ l of 1% phosphotungstic acid in 0.5M HCl, followed by centrifugation at 10,000 \times g for 5 min. The pellets were washed once with the same phosphotungstic acid solution and once with methanol, dissolved in 0.5 ml of 0.05M NaOH and 1M NaBH₄, and then incubated 30 h at 45°C.

At the end of the incubation period, the solution was neutralized with acetic acid to pH 6 and then lyophilized. The dehydrated products were dissolved in 50 μ l of water, and then desalted by gel filtration on a Sephadex G-15 column (0.5 × 5 cm) equilibrated and eluted with water. The radioactive fractions were subjected to thin layer chromatography for identification of the products without further purification. NeuAco2,6GalNAc-ol and GlcNAc β 1,3[NeuAco2,6]GalNAc-ol from native BSM in two different developing solvent were co-migrated. The ratio of the transferred sialyl residue was 1:0.9:0.6. The results of the co-migration of Sialylated GalNAc-SerNAc with with NeuAco2,6GalNAc-SerNAc in the two different solvent systems indicate that the protein SB-690 of the present invention forms the NeuAco2,6 linkage to GalNAc that is directly attached to Ser or Thr residues in glycoproteins.

Media from cells transfected with pcDSB-690 contained sialyltransferase activity and it provide strong evidence that the protein SB-690 of the present invention expressed by pcDSB-690 was secreted out of cells while retaining sialyltransferase activity. On the other hand, media obtained from cells transfected with cDSB-BGL had no sialyltransferase activity.

The acceptor specificity of the protein SB-690 of the present invention was examined with the concentrated COS-7 cell culture medium transfected with pcDSB-690. As shown in Table 1, asialo-mucin, fetuin and asialo-fetuin served as good acceptors. Remarkably, fetuin was shown to be a better acceptor than asialo-fetuin (Baubichon-Cortay, H. et al., Carbohydr. Res., 149, 209-223, 1986; and Brockhausen, I. et al., Biochemistry, 29, 10206-10212, 1990). Other glycoproteins, oligosaccharides and glycolipids did not serve as acceptors, except GalNAc-SerNAc. These data suggest that the acceptor site is GalNAc directly attached to Ser or Thr residues in glycoproteins through an α-glycoside linkage.

Table 1

Acceptor specificity of the protein	SB-690 of the invention
Acceptor	Pmoles/hr/10 µl medium
Fetuin	142
Asialo-fetuin	96
α1 acid glycoprotein	. 6
Asialo-α1 acid glycoprotein	4
Bovine submaxillary mucin	15
Bovine submaxillary asialo-mucin	186
Ovomucoid	7
Asialo-ovomucoid	0
Galβ1,3GlcNAcβ1,3Galβ1,4Glc	۰ 0
Galβ1,4GlcNAc	0
Galβ1,3GalNAc .	0
GalNAcβ1,4 Gal	0
Galβ1,4Glc	0
Galactose	0
Ganglioside mixture	0
Ganglioside GD1a	0
GalNAc-SerNAc	4
Benzyl-GalNAc	2

^{*} A number of 0 indicates lass than 1 pmol/hr/10 µl medium.

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So far cloned sialyltransferases only exhibit acceptor specificity for the Gal-moiety. While the GalNAca2,6-sialyl-transferase P-B1 and protein SB-690 of the present invention exhibit acceptor specificity for the GalNAc-but not the Gal-moiety. The following evidence supports that GalNAca2,6-sialyltransferase P-B1 and the protein SB-690 of the present invention have the activity of GalNAca2,6-sialyltransferase, which transfer CMP-NeuAc with an a2,6-linkage onto a GalNAc residue O-linked to Thr/Ser of a glycoprotein:

- (i) The expression of pcDSB-690 in COS cells reveals the remarkable acceptor specificity for only the GalNAc moiety bound to Ser/Thr residues, while no detectable enzyme activity was found toward the other substrates tested (Table 1).
- (ii) The sialylated products obtained from bovine submaxillary asialo-mucin and GalNAc-SerNAc were shown to have sialic acid bound to the GalNAc moiety through an α2,6-linkage.

The two types, i.e., bovine submaxillary gland- and liver (brain)- types, of GalNAc-α6ST were reported, which have the different acceptor specificity (Bergh, M.E. et al., J. Biol. Chem., 258, 7430-7436, 1983). The former enzyme has the broad specificity toward GalNAc, Galβ1,3GalNAc and NeuAco2,3Galβ1,3GalNAc, whereas the latter has only toward NeuAco2,3Galβ1,3GalNAc moiety of glycoproteins. The acceptor specificities of the GalNAco2,6-sialyltransferase P-B1 and the protein SB-690 of the present invention were found to be similar to that of the former enzyme.

Examination of the acceptor site of asialo-mucin showed that NeuAco2,6GalNAc-Ser/Thr was the most abundant product. However, considering the ratio of glycoconjugates in bovine submaxillary asialo-mucin, i.e., GalNAc-Ser/Thr, GlcNAcβ1,3GalNAc-Ser/Thr, and Galβ1,3GalNAc-Ser/Thr amounted to 65%, 25%, and 5%, respectively (Tsuji, T. and Osawa, T., Carbohydr. Res., 151, pp.391-402, 1986), GalNAco2,6-sialyltransferase P-B1 and the protein SB-690 of the present invention seem to have the following acceptor preference: Galβ1,3GalNAc-Ser/Thr > GlcNAcβ1,3GalNAc-Ser/Thr. On the other hand, the facts that almost all the radioactivity was released on weak alkali treatment and that fetuin is preferred over asialo-fetuin (Table 1) indicate that NeuAco2,3Galβ1,3GalNAc-Ser/Thr is a preferred substrate over Galβ1,3GalNAα-Ser/Thr, as reported for calf liver (Bergh, M.E. et al., J. Biol. Chem., 258, 7430-7436, 1983) and rat brain (Baubichon-Cortay, H. et al., Carbohydr. Res., 149, 209-223, 1986) GalNAco2,6-sialyl-transferases.

The sialylation of GalNAc-SerNAc was much slower than that of corresponding residues on asialo-mucin (Table 1). Brockhausen et al. (Brockhausen et al., Biochemistry, 29, 10206-10212, 1990) showed that a length of at least five amino acid is required for efficient synthetase activity. A similar effect of the peptide portion directly on GalNAco2,6-sialyltransferase P-B1 and the protein SB-690 of the present invention is also suggested from this observation (Table 1).

The regents and the like used in the above preparation examples (A) and (B) were as follows: Fetuin, asialo-fetuin, bovine submaxillary mucin, α1-acid glycoprotein, galactose β1,4-N-acetylgalactosamine, CMP-NeuAc, lacto-N-tetraose, benzyl-GalNAc, N-acetyllactosamine, and Triton CF-54 were obtained from Sigma (St. Louis, USA). CMP[¹⁴C]NeuAc(11 GBq/mmole) was obtained from Amersham (U.K.). N-Acetylgalactosamine β1,4-galactose was a gift from Dr. Kajimoto (The institute of Physical and Chemical Research, RIKEN, Wako-shi, Saitama-ken, Japan). 2-Acetamide and 2-deoxygalactosyl-αN-acetylserine (GalNAc-SerNAc) were synthesized according to Grundler and Schmidt (Grundler G., and Schmidt R.R., Liebigs Ann. Chem., 1984, 1826-1847, 1984). NeuAcα2,6-GalNAc-SerNAc was prepared from NeuAcα2,6GalNAc-Ser (MECT) by acetylation with anhydroacetate in pyridine-water. NeuAcα2,6GalNAc-ol and GlcNAcβ1,3[NeuAcα2,6]GalNAc-ol were prepared from bovine submaxillary mucin according to Tsuji and Osawa (Tsuji, T. and Osawa T., Carbohydr. Res., 151, 391-402, 1986) and identified by 270MHz ¹H and ¹³C NMR (Savage, A.V. et al., Eur. J. Biochem., 192, pp. 427-432, 1990; and Savage, A.V. et al., Eur. J. Biochem., 193, 837-843, 1990). Synthetic primers were synthesized with the Applied Biosystem 394 DNA synthesizer. Restriction endonucleases Smal, EcoRl, BamHl, Hindill, Sacl, Xhol, Bglll and Pstl were from Takara (Japan).

(C) Preparation of GalNAco2.6-sialyttransferase P-B3

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In order to obtain cDNA clones of GalNAco2,6-sialyltransferases, PCR with two degenerate oligonucleotides (ST-107 and ST-205) was performed with chick embryo cDNA as a template. The fragment of the desired size of approximately 150 bp was purified by agarose gel electrophoresis. As a result of sequencing of the PCR products, it was revealed that they included those encoding Galβ1,4GlcNAco2,6-sialyltransferase (Kurosawa, N., et al., Eur. J. Biochem., 219, 375-381, 1994) and GalNAco2,6-sialyltransferase P-B1, as well as a PCR product encoding a novel amino acid sequence, pCRB3. The identity of the sialylmotif of pCRB3 with those of above-mentioned sialyltransferases was 65 through 57%.

In order to identify the complete coding sequence of the gene, a young chicken testis cDNA library was screened with the cDNA insert of pCRB3. The screening about 5×10^5 independent clones yielded one positive clone, λ CEB3-T20, which has an insert size of 2.05 kb.

The nucleotide sequence of the cDNA done included an open reading flame of 1212 bp, coding for 404 amino acids with a molecular mass of 45.8 kDa. The open reading frame starts with a methionine codon at nucleotide 1, with

a conventional translation initiation sequence (Kozak, M. Nature, 308, 241-246, 1984), and ends with a TGA stop codon at nucleotide 1213. The open reading flame is flanked by a 5'-untranslated sequence of 384 bp and a 3'-untranslated sequence of 451 bp. The DNA sequence 5' of the initiation site contains stop codons in all three reading frames. The nucleotide sequence and deduced amino acid sequences of λ CEB3-T20 are shown in the SEQ ID No.3 of the sequence listings. The GalNAco2,6-sialyltransferase having this amino acid sequence was designated as P-B3.

This GalNAco2,6-sialyltransferase P-B3 (when the GalNAco2,6-sialyltransferase P-B1 is referred to as ST6GalNAcA, this enzyme is occasionally referred to as ST6GalNAcB) has type II transmembrane domain, containing a 17-amino acid N-terminal hydrophobic sequence bordered by charged residues, as has been found for all sialyltransferases cloned to date. Comparison of the primary sequence of GalNAco2,6-sialyltransferase P-B3 with other amino acid sequences in DNA and protein data banks revealed similarities in two regions to all of the cloned sialyltransferases.

One region (sialylmotif L) in the center of the GalNAcα2,6-sialyltransferase P-B3, consisting of a 45 amino acid stretch, shows 64-24% sequence identity, whereas the other, in the COOH-terminal portion (sialylmotif S, residues 333-355), exhibits 78-43% identity. The overall amino acid sequence identity of GalNAcα2,6-sialyltransferase P-B3 is 10% to chick Galβ1,4GlcNAα2,6-sialyltransferase (Kurosawa, N., et al., Eur. J. Biochem., 219, 375-381, 1994), 13% to chick Galβ1,3GalNAcα2,3-sialyltransferase (Kurosawa N. et al., Biochem. Biophys. Acta., 1244, 216-222, 1995), and 32% to chick ST6GalNAcA (22), respectively. These results suggest that the cloned gene belongs to the sialyltransferase gene family.

Details of the experiments are as follows.

Polymerase chain reaction (PCR)

PCR was performed using degenerate primers [5' primer ST107: TGGGCCTTGGII(A/C)AGGTGTGCTGTTQ, and 3' primer ST-205: AGGCGAATGGTAGTTTTTG(A/T)GCCCACATC] deduced from conserved regions in Galβ4GicNAc-α6STRL (Weinstein, J. et al., J. Biol. Chem., 262, 17735-17743, 1987), Galβ4GicNAc-α6STHP (Grundmann, U. et al., Nucleic acids Res., 18, 667, 1990), and Galβ3GalNAc-α3STPS (Gillespie, W. et al., J. Biol. Chem., 267, 21004-21010, 1992). To obtain cDNA, poly(A)-rich RNA (2 μg) from 3 day-old chick embryos was incubated with an oligo-dT primer (Pharmacia), 1 mM each of dATP, dCTP, dGTP and dTTP, and 2 U/μl of RNase inhibitor (Promega) in 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.5 mM MgCl₂ and 0.001% gelatin in 50 μl for 10 min at 0 °C, and then for additional 60 min at 42°C after the addition of 100 μU Moloney murine leukemia virus reverse transcriptase (BRL).

After heating at 94 °C for 3 min, cDNA prepared from 0.2 μ g of poly(A)-rich RNA was used for the PCR experiment in a mixture comprising 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.25 mM MgCl₂, 0.001% gelatin, 200 μ M of each dATP, dCTP, dGTP and dTTP, 2U of Taq DNA polymerase (Promega), and 40 pmoles of each PCR primer in 50 μ l. PCR amplification, 35 cycles, was carried out, each cycle consisting of denaturation at 96°C for 45 sec, annealing at 50 °C for 60 sec, and extension at 72°C for 60 sec. The PCR products were developed on a 3% agarose gel. The DNA fragment corresponding to 150 bp was eluted from the gel (Qiaex kit; Qiagen), blunt-ended, kinated, and then subcloned into the Smal site of pUC119, and finally sequenced.

Construction of a cDNA library

Total RNA was prepared from chick embryos (6 day-old) by the guanidinium thiocyanate method, followed by centrifugation in a 5.7 M CsCl solution (Sambrook, J., Molecular Cloning: a Laboratory Manual, 2nd edition). Poly(A)rich RNA was purified with Oligotex-dT30 (Takara), and then employed for the construction of a cDNA library using λZAPII (Stratagene) and cDNA synthesis kits (Pharmacia) with an oligo-dT primer and random primers.

45 Screening of the cDNA library

The amplified cDNA library (1 \times 10⁶ plaques) was screened with the chick embryo PCR fragments. The plaque-transferred filters were hybridized with ³²P-radiolabeled DNA probes for 12 h at 65°C in 5 \times SSC, 0.2% SDS, 5 \times Denhardt's solution, and 10 μ g/ml denatured salmon sperm DNA. The filters were then washed twice at 65°C for 20 min in 2 \times SSC, 0.1% SDS. To obtain plasmids from the isolated phage clones, phagemid rescue was performed according to the instructions of the manufacturer of the λ ZAPII cloning kit (Stratagene). cDNA inserts were excised directly as Bluescript plasmids. Plasmids were produced by the standard molecular cloning method according to Sambrook, et al. (Sambrook, J. et al., Molecular Cloning: a Laboratory Manual, 2nd ed.)

DNA sequence analysis

The DNA sequences of the inserts were determined by the dideoxy-chain termination method (Sanger, F. et al., Proc. Natl. Acad. Sci. USA, 74, 5463-5467, 1977) using single-strand DNA as a template for T7-DNA polymerase. The sequencing reaction and electrophoresis were carried out using an AutoRead DNA sequencing kit and a DNA

sequencer (Pharmacia). Single Strand DNA was prepared from Escherichia coli XL-Blue (Stratagene) after superinfection with helper phage R408 (Stratagene). The sequence data were analyzed with a computer using PC/Gene (Teijin System Technology).

To confirm the existence of the gene, Southern blot analysis was performed for chicken genomic DNA. Hybridization of the cDNA insert of pCRB3 for chicken genomic DNA gave a single band on digestion with EcoRI and two bands with BamHI. This simple hybridization pattern indicates that the cloned cDNA was a single copy gene. Southern blot analysis of genomic DNA from mouse and monkey with the pCRB3 probe under low stringency conditions suggested that this gene is conserved across species. For Southern blot, each 7.5 µg of genomic DNA prepared from mouse brain, COS-7 cells and chicken testes were digested with restriction enzyme and then size-fractioned on 0.6% agarose gels.

The mRNA size and distribution of the GalNAco2,6-sialyltransferase P-B3 gene were determined by Northern blot analysis. Analysis of RNA from 3, 6, 8, 10 and 12-day old embryos revealed two RNA species of 4.5 kb and 2.2 kb. The 4.5 kb mRNA was expressed abundantly at all embryonic stages examined, while not expressed in adult tissues. The less abundant 2.2 kb mRNA was expressed at the early embryonic stage, being abundant at the late embryonic stage and in adult tissues. The size of the 2.2-kb transcript suggests that the obtained cDNA clone (λCEB3-T20) was close to full length. For Northern blots, 5 μg of poly(A)-rich RNAs from chick embryo and 10 μg of all RNA from chicken tissues were size-fractioned on formaldehyde-agarose gels.

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Sialyltransferases previously known exhibit remarkable tissue-specific expression, which is considered to be correlated with the existence of cell type-specific carbohydrate structures (Paulson, J.C. and Colley, K.J., J. Biol. Chem., 264, pp.17615-17618, 1989). The results of Northern blotting indicate that the pattern of expression of sialyltransferase P-B3 changes. The precise structure of embryo-specific 4.5 kb mRNA has not been known. However, the production of two different sizes of mRNAs from the sialyltransferase e P-B3 gene suggests that they are very likely to be generated through alternative splicing and alternative promoter utilization mechanisms as observed for Galβ1,4GicNAc-α2,6-sialyltransferase (Galβ4GicNAc-α6STRL) and Galβ1,3(4)GicNAc-α2,3-sialyltransferase (Galβ3(4)GicNAc-α3STRL) (Weinstein, J. et al., J. Biol. Chem., 262, 17735-17743, 1987; and Wen, D.X. et al., J. Biol. Chem., 267, 21011-21019, 1992). This hypothesis is supported by the results of Southern hybridization, which showed the existence of a single copy gene for sialyltransferase P-B3.

A 1.3 kb DNA fragment encoding the full length sialyltransferase P-B3 was amplified using synthetic oligonucleotide primers (5'-ACGGCG<u>CTCGAG</u>CCAACCCGGAGAGCAGCG-3', and 5'-CGTTGC <u>CTCGAG</u>AGTCCTTGCAGT-GGGACT-3', synthetic Xhol site underlined). The amplified DNA fragment was digested with Xhol and inserted into the Xhol site of the expression vector pcDSRα (Takebe, Y., Mol. Cell. Biol., 8, pp.466-472, 1988) to yield recombinant plasmid pcDB3ST. The insert of the plasmid was sequenced to confirm the absence of possible polymerase chain reaction errors.

COS-7 cells were transfected with 5 µg of the recombinant plasmid pcDB3ST using the DEAE-dextran method (McCutchan, J.H. and Pagano, J.S., J. Natl, Cancer Inst., 41, 351-357, 1968).

After 48 h of the transfection, the cultured cells (1 × 10⁷) were harvested, washed with phosphate-buffered saline, and then resuspended in 2 ml of buffer comprising 20 mM MnCl₂ and 25 mM MES, pH 6.0. The cell suspension was centrifuged at 30,000 × g for 30 min, the cell pellet was resuspended in 0.5 ml or 1% Triton X-100, 50 mM NaCl, 5 mM MnCl₂, 25 mM MES, pH 6.0, and then subjected to sonication. After centrifugation at 30,000 × g for 30 min, the supernatant was concentrated 10-fold on Centricon 30 filters (Amicon), and then used for following assays.

The enzyme assays with glycoproteins, oligosaccharides and glycolipids as acceptors were performed in the presence of 0.1 M sodium cacodylate buffer (pH 6.0), 10 mM MgCl₂, 0.5% Triton CF54, 12 μ M CMP-[¹⁴C]NeuAc (1.5 kBq), 1 mg/ml acceptor substrate, and 1 μ l of COS cell lysate (in a final volume of 10 μ l), with incubation at 37 °C for 1 hr. At the end of the incubation period, the reaction mixtures were subjected to SDS-PAGE for glycoproteins as acceptors, or were subjected to chromatography on HPTLC plates (Merck, Darmstadt, Germany) with a solvent system of ethanol/1-butanol/pyridine/acetic acid/water (100:10:10:3:30) for oligosaccharides and glycolipids as acceptors. Sialylated acceptors were quantified with a BAS2000 radio image analyzer (Fuji Photo Film, Japan).

Identifications of sialylated products were as follows. Reduced oligosaccharides were obtained from resialylated glycoproteins by β-elimination as described by Carlson (Carlson, D.M., J. Biol. Chem., 243, pp616-626, 1968). AsialoBSM was sialylated with CMP-[¹⁴C]NeuAc in a pcDB3ST-transfected COS-7 cell lysate under the same conditions as above. The radiolabeled oligosaccharides released from fetuin were digested with NDV sialidase, and then subjected to thin layer chromatography for identification of the products without further purification. Oligosaccharides released from BSM were used as standards. AsialoBSM and asialofetuin were [¹⁴C]-sialylated with the GalNAco2,6-sialyltransferase P-B1 and Galβ1,3GalNAco2,3-sialyltransferase (Lee, Y.-C., et al., Eur. J. Biochem., 216, pp. 377-385, 1993), respectively, and the oligosaccharides were prepared by β-elimination. The resulting [¹⁴C]NeuAco2,6GalNAcol, Galβ1,3([¹⁴C]NeuAco2,6)GalNAcol and [¹⁴C]NeuAco2,3Galβ1,3GalNAcol were used as radio-labeled standards.

When fetuin was used as the acceptor, the acceptor was only sialylated by the lysate of COS-7 cells transfected with pcDB3ST. The expressed GalNAco2,6-sialyltransferase P-B3 exhibited strong activity toward fetuin and asialofetuin, and weak activity toward asialoBSM, whereas no significant activity was observed toward BSM or other glycopro-

teins having only N-glycosidically linked oligosaccharides (e.g., α 1-acid glycoprotein, ovomucoid, asialo- α 1 acid glycoprotein and asialo-ovomucoid) (Table 2).

In addition, oligosaccharides or glycosphingolipids could not serve as acceptors for the GalNAc α 2,6-sialytransferase P-B3 of the present invention. [14 C]NeuAc residues incorporated into fetuin by the enzyme were resistant to treatment with N-glycanase or NDV sialidase. The radiolabelled oligosaccharides released from fetuin were comigrated with Gal β 1,3(NeuAc α 2,6)GalNAc-ol after treatment with NDV sialidase. These results indicate that sialic acid residues were transferred through α 2,6-linkages on GalNAc residues of O-glycosidically linked oligosaccharides of fetuin. Thus, the expressed enzyme apparently has GalNAc α 2,6-sialytransferase activity. However, asialoBSM was a much poorer acceptor than fetuin and asialofetuin for this GalNAc α 2,6-sialytransferase P-B3 of the present invention. The acceptor substrate specificity is different from that of the GalNAc α 2,6-sialytransferase P-B1 for which asialoBSM serves as a much better acceptor than asialofetuin.

To define the substrate specificity of the GalNAco2,6-sialyltransferase P-B3 of the present invention, fetuin was sequentially treated with sialidase (Vibrio cholerae) and β-galactosidase (bovine testes), and the resulting asialofetuin and agalacto-asialofetuin were used as acceptors. The incorporation of NeuAc-residues for the sialidase-treated fetuin was increased 1.5-fold of that for native fetuin. Three O-glycosidically linked oligosaccharides are known to be contained in fetuin, two of which are NeuAco2,3Galβ1,3GalNAc and the other is NeuAco2,3Galβ1,3(NeuAco2,6)-GalNAc (Spiro, R.G. and Bhoyroo, V. D., J. Biol. Chem., 249, 5704-5717, 1974). Accordingly, GalNAc residues in two of the three O-linked oligosaccharides can serve as acceptors in native fetuin, whereas those in all O-linked oligosaccharides in asialofetuin can be sialylated by the GalNAco2,6-sialyltransferase P-B3 of the present invention.

Furthermore, agalacto-asialofetuin could not serve as an acceptor of the GalNAc α 2,6-sialyltransferase P-B3 of the present invention, and only Gal β 1,3([14 C]NeuAc α 2,6)GalNAc-ol, but not [14 C]NeuAc α 2,6-GalNAc-ol, was detected for the oligosaccharides released from asialoBSM incubated with the enzyme by β -elimination.

The characteristics of the GalNAco2,6-sialyltransferase P-B3 of the present invention revealed by the above experiments can be summarized as follows:

- (1-i) Fetuin and asialofetuin, which contain the O-glycosidically linked (NeuAcα2,3)Galβ1,3GalNAc sequence (Spiro, R.G. and Bhoyroo, V.D., J. Bio. Chem., 249, 5704-5717, 1974), served as good acceptors, but asialoBSM, in which only 5% of the total carbohydrate chains contain Galβ1,3GalNAc sequences (Tsuji, T. and Osawa, T., Carbohydr. Res., 151, 391-402, 1986), served as a much poorer acceptor; and
- (1-ii) the protein portion is essential for the activity of this sialyltransferase, since Gal β 1,3GalNAc α 1-Bz as well as asialoGM1 (Gal β 1,3GalNAc β 1,4Gal β 1,3Glc β 1-Cer) and GM1b (NeuAc α 2,3Gal β 1,3GalNAc β 1,4Gal β 1,3Glc β 1-Cer) did not serve as acceptors.
- (2) This sialyltransferase did not exhibit activity toward asialofetuin treated with β-galactosidase (agalacto-asialofetuin).
- (3) Only Galβ1,3([¹⁴C]NeuAc-α2,6)GalNAc-ol was detected in the oligosaccharides released from [¹⁴C]sialylated asialoBSM although about 60% of the carbohydrate chains of asialoBSM are GalNAc-O-Ser/Thr (Tsuji, T. and Osawa, T., Carbohydr. Res., 151, 391-402, 1986).

These results clearly suggest that the acceptor substrate of the enzyme of the present invention having catalytic activity, i.e., transfer of CMP-NeuAc with an α2,6-linkage onto a GalNAc residue O-linked to Thr/Ser of a glycoprotein, requires Galβ1,3 GalNAc sequence of O-glycoside linked oligosaccharide, whereas α2,3 linkage-sialic acid residues linked to galactose residues are not essential for the activity. Therefore, the enzyme P-B3 first cloned by the present invention is a novel type of GalNAcα2,6-sialyttransferase. The primary sequence of GalNAcα2,6-sialyttransferase P-B3 from the 45 amino acid regions at the molecular center (sialylmotif L) to the COOH-terminal (residues: 180-404) exhibits high sequence homology to that of GalNAcα2,6-sialyttransferase P-B1 (Fig. 4: the identity is 48%). The conserved regions unique to these GalNAcα2,6-sialyttransferases may be correlated with their enzymatic function of transferring sialic acid to the GalNAc-moiety via an α2,6-linkage.

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Table 2

Acceptor substrate specificity ferase P-B3 of	
Acceptor	Specificity
	pmol/h/µl enzyme fraction
Fetuin	28
Asialofetuin	35
BSM	0.5
AsialoBSM	5.2
α1-Acid glycoprotein	0
Asialo-α1-acid glycoprotein	1.2
Ovomucoid	0
Asialo-ovomucoid	1.0
Galβ1,3GalNAcα1-Bz	0
GaiNAca1-Bz	0
GalNAc-SerNAc	0
AsialoGM1	. 0
GM1b	0
Ganglioside Mixture	0
0 indicates less than 0.5 pmo	Vh.

The regents, samples and the like used in the above preparation example (C) were as follows. Fetuin, asialofetuin, bovine submaxillary mucin, α1-acid glycoprotein, galactose β1,4-N-acetylgalactosamine, CMP-NeuAc, Galβ1,3GalNAcα1-Bz, GalNAcα1-Bz and Triton CF-54 were obtained from Sigma (St. Louis, USA). CMP-[¹⁴C]NeuAc (11 GBq/mmole) was obtained from Amersham (U.K.). 2-Acetamide and 2-deoxygalactosylαN-acetylserine (GalNAc-SerNAc) was synthesized according to Grundler and Schmidt (Grundler G., and Schmidt R.R., Liebigs Ann. Chem., 1984, 1826-1847, 1984). NDV-sialidase and sialidase from Vibrio cholerae were purchased from Oxford Glycosystems (U.K.) and Boehringer Mannheim (Germany), respectively. p-Galactosidase from bovine testes was obtained from Boehringer Mannheim (Germany). Synthetic primers were synthesized with the Applied Biosystem 394 DNA synthesizer. Restriction endonucleases were obtained from Takara (Japan)

(D) Purification of sialyttransferase expressed in microorganisms

Plasmid construction

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An initiation codon and cloning sites were attached by PCR to mouse Galβ1,4GlcNAco2,6-Sialyltransferase cDNA (Hamamoto, T. et al., Bioorg. Medicin. Chem., 1, 141-145, 1993). 5'-TGGCATATGGGGAGCG ACTATGAGGCTCT-3'containing an Ndel site was used as a sense primer and 5'-ATGAGGATCCCTGGCTCAACAGCG-3' containing a BamHI site as an antisense primer. The resulting PCR fragment (1152 bp) contained the initiation codon and a region coding for a polypeptide from the 29th amino acid residue to the C-terminal end of the enzyme, and lacked the cytosolic and transmembrane domains. The fragment was incorporated into expression vector pET3b (Studier, F.W. et al., Method. Enzymol., 185, 60-89, 1990) at the Ndel-BamHI site (located downstream of the T7 promoter). The resulting recombinant vector was named as pET3-MBS. The nucleotide sequence of the PCR fragment is shown as the SEQ ID No.4 in the sequence listings.

Enzyme expression

E. coli JM109(DE3) cells transfected with the vector pET3-MBS were cultured in 100 ml LB medium supplemented with 100 μg/ml ampicillin at 37 °C. When the optical density at 600 nm reached 0.2-0.4, production of the recombinant protein was initiated with induction of T7 RNA polymerase by the addition of 2 mM IPTG (isopropylβ-D-thiogalatopyranoside). The recombinant enzyme, lacking the cytosolic and the transmembrane domain, was accumulated in the form of insoluble inclusion bodies in the cells. The growth rate of the JM109(DE3) cells transfected with pET3-MBS was the same as that of the non-transfected JM109(DE3) cells both on agar plates and in liquid culture. After 2 h cultivation, the cells were harvested (ca. 1 g wet weight), suspended in 10 ml of 20 mM Tris-HCI (pH 8.0), and then treated with lysozyme (0.1 mg/ml) and DNase I (0.01 mg/ml) for 30 min. Triton X-100 was added to a final concentration of 1%, and insoluble fraction was collected by centrifugation at 12,000 x g for 15 min at 4 °C. The precipitate was suspended in 3 ml of 10 mM Tris-HCI (pH 7.4) and stored at -30°C before use.

Solubilization and renaturation

To 0.5 ml of the above suspension, 0.48 g solid urea, 60 μ of 5 M NaCl, 20 μ l of 1 M Tris-HCl (pH 7.4) and water were added to final volume of 1 ml (final concentration: 8 M urea, 0.3 M NaCl; 20 mM Tris-HCl, pH 7.4). The precipitate was extracted for 30 min at 10°C, followed by centrifugation at 12,000 \times g for 15 min. Most of the extracted protein had the molecular mass of 42k dalton. Where 5.7 M urea buffer was used for the extraction, 80% of the enzyme was recovered.

The 0.1 ml aliquots of extract containing 8 M urea were diluted with each 1.9 ml of a renaturation composition (standard composition: 2 M urea, 0.5 M NaCl, 10 mM lactose, 0.5 mM EDTA, and 20 mM MOPS-NaOH, pH 7.0) to a final protein concentration of about 0.02 mg/ml. The solution was left at 40°C for 12 h, and then diluted again with an equal volume of the renaturation composition, thereby reducing the urea concentration to half (approximately 1.2 M), and then the mixture was left at 40 °C for additional 48 h. Then, sialyltransferase activity was measured to analyze the effects of the components of the renaturation composition at this point (Table 3). The resulting enzymes were further dialyzed against the renaturation composition to remove residual urea and the reducing agents over 48 h at 4 °C. The samples were concentrated approximately 20 times with Centricon-30 (Amicon).

Sialyttransferase assay

The activity of the sialyltransferase was measured with 50 μM CMP-[¹⁴C]NeuAc (0.9 Bq/pmole) as a donor substrate, and 5 mM Galβ1,4GlcNAc (N-acetyllactosamine) as an acceptor substrate. Reaction mixture was added with 1 mg/ml bovine serum albumin, 1 μl of the enzyme solution, and 50 mM sodium cacodylate (pH 6.0) to a total volume of 10 μl, and incubation was continued at 37 °C for 1 h. Then, the samples were applied to silica gel60 HPTLC plate (Merck Germany) and developed with ethanol/pyridine/n-butanol/acetic acid/water (100:10:10:3:30) as a developing solvent. The radioactivity transferred on each plate was determined with a radio image analyzer BAS2000 (Fuji Photo Film, Japan, Lee, Y.-C. et al., Eur. J. Biochem., 216, 377-385, 1993). One unit of enzymatic activity was defined as an amount catalyzing 1 μmole of sialic acid transfer per minute. The acceptor preference as to oligosaccharide branches was examined using a N-acetyllactosamine type biantennary pyridylamino-oligosaccharide as an acceptor substrate and analyzed fluorophotometrically by HPLC.

When the 8 M urea extract was dialyzed without dilution at 4 °C, almost no activity of the enzyme precipitated at concentration of less than 0.5 M was recovered. The results of the optimum dilution conditions at 48 h after the second dilution are shown in Table 3 set out below. In the table, the standard renaturation composition was comprised of: 2 M urea, 20 mM Tris-HCl, 0.3 M NaCl, 20 mM lactose, and 0.5 mM EDTA (pH 7.4), and as to other compositions, deviations from the standard composition are indicated.

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Table 3

The effects of various con Galβ1,4GalNAco2,	
Renaturation conditions	Relative activity compared to standard
Standard composition	1
pH.9.5, Tris-HCl 20 mM	0* .
pH 8.0, Tris-HCl 20 mM	0.6
pH 7.0, MOPS-NaOH 20 mM	2.5
pH 6.0, MES-NaOH 20 mM	1.5
0.5 M NaCl	2
0.1 M NaCl	0.2
0.01 M NaCl	0
0 mM lactose	0.5
1 M. urea	1.5
0 M urea	0.6

^{*} A value of 0 indicates less than 5% of the control.

The maximum renaturation was observed with 0.5 M NaCl (pH 7.0) in the standard composition, and these compositions were used in further experiments. After three independent renaturation experiments were carried out under this condition, total recovered activities were 0.4-0.8 mU/0.1 ml extract. The enzymes at this stage of renaturation showed high Km values for CMP-NeuAc and N acetyllactosamine, 0.14 mM and 20 mM, respectively. Under the conditions tested, reducing agents (DTT and β-mercaptoethanol) inhibited the enzyme activity, which may due to carryover of urea at the concentration of 0.1 M in the assay mixture. In addition, very little activity was observed at 12 h after the second dilution, which apparently indicates that a refolding process of the polypeptide is very slow at the test temperature. Almost the same activity, as that in the process without the use of the reducing reagents, was obtained by the following process: the 8 M urea extract was diluted with 20 volumes of the renaturation composition containing 2 M urea, 20 mM MOPS-NaOH, pH 7.0, 0.5 M NaCl, 20 mM lactose, and 0.5 mM EDTA in the presence of 1 μM or 1 mM reducing regents, and then samples were left at 4 °C for 12 h and diluted to reduce the urea concentration to half, and the residual urea and reducing reagents were removed by dialysis. The results are shown in Table 4.

Table 4

Reducing regent	Specific activity (mU/mg)
None	7
1 μM DTT	6
1 mM DTT	12

The substrate specificity of renatured mouse Galβ1,4GlcNAco2,6-sialyltransferase was assayed using each 2 mg/ml of substrates. The products were analyzed by HPTLC. HPTLC was performed using ethanol/pyridine/n-buta-nol/acetic acid/water (100:10:10:3:30) as a developing solvent when oligosaccharides and glycoproteins were used as acceptors, and chloroform:methanol:0.5% CaCl₂ (55:45:8) as a developing solvent when glycolipids were used as acceptors. The substrate specificity and kinetic parameters of the renatured enzymes were similar to those of the enzyme obtained from rat liver.

The results are shown in Table 5 and Table 6.

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Table 5

Substrate Relative Activity to Gal61,4GlcNAc Renatured mouse Rat liver Galß1,4GlcNAco2,6-sia-Galβ1,4GlcNAcαa2,6-sialyltransferase lyltransferase **Fetuin** 0* 0.25 Asialofetuin 1.5 0.97 a1 acid glycoprotein 0.1 0.1 Asialo-a1 acid glycoprotein 2.1 1 Bovine submaxillary mucin 0 0 Bovine submaxillary asialo-mucin 0 Ó Lacto N-tetraose 0 0 Galβ1,4GlcNAc Gal_B1,3GlcNAc 0 0 GalNAc_{β1,4}Gal n 0 Galp1,4Glc 0 0 Gal 0

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Table 6

Substrate	Km	(mM)
	Renatured mouse Galβ1,4GicNAco2,6-sia- lytransferase	Rat liver Galβ1,4GlcNAcαa2,6-sia- lyttransferase
CMP-NeuAc*	0.08	0.04
N-acetyllactosamine	6.5	5
Asialo-orosomucoid**	0.4	0.2

^{*} Measured with N-acetyllactosamine as the acceptor.

Galβ1,4GlcNAco2,6-sialyltransferase is capable of recognizing the different branches of biantennary glycopeptides of the N-acetyllactosamine type (Joziasse, D.H. et al., J. Biol. Chem., 260, 714-719, 1985; and Van den Eijnden D.H. et al., Biochem. Biophys. Res. Comm., 92, 839-845, 1980). A desialylated biantennary PA-oligosaccharide was sialylated by the enzyme renatured according to the method of the present invention and then analyzed with HPLC. The assays were performed using 10 pmoles of acceptor substrates and 0.1 mM CMP-NeuAc in a final volume of 5 μl. The reaction mixtures were incubated at 37°C for 1 h, and the reaction was stopped by the addition of 90 μl of cold water. To identify sialylated pyridylamino oligosaccharides, each reaction mixture was subjected to HPLC analyses equipped with a reversed-phase column (Shimpack CLC-ODS, 0.6 cm × 15 cm, Shimazu, Japan). The column was equilibrated with mixture of 70% solvent A (10 mM sodium phosphate, pH 3.8) and 30% solvent B (0.5% n-butanol, 10 mM sodium phosphate, pH 3.8), and eluted at the flow rate of 1 ml/min with a linear gradient of solvent B to 60% over 30 min at 55°C. Pyridylamino oligosaccharides were detected fluorophotometrically (excitation at 320 nm and emission at 400 nm), and

^{*} A value of 0 indicates less than 2% of the control.

[&]quot;Concentration expressed as terminal galactose residues.

the results indicated that the renatured enzyme showed higher preference for galactose residues on Man α 1,3 branches rather than for galactose residues on Man α 1,6 branches like the native enzyme.

By competely remove urea, the renatured enzyme restored its resistance to reducing agents. In addition, more than 10 times activation was recovered by renaturing with the addition of divalent cations. While not bound by any specific theory, where dialysis is carried out for a prolonged period of time against a dialysis buffer containing 0.5 mM EDTA in the presence of urea, divalent cations, which are tightly bound to the enzyme to maintain the proper conformation of the enzyme, may be lost. Where the enzyme was renatured in the renaturation composition containing 1.2 M urea, the addition of divalent cations increased the activity. The results obtained are shown in Table 7. In the table, the activities are shown as relative values to that obtained by no addition of reagents. The specific activity of the renatured enzyme was 0.15 U/mg protein when measured with 5 mM MnCl₂, which is about 2% of that of the enzyme obtained from rat liver (weinstein, J. et al., J. Biol. Chem., 257, pp.13835-13844, 1982). The overall recovery of the enzyme was 0.1 U/100 ml culture medium.

Table 7

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Reagent	Renatured mouse Galβ1,4GlcNAcα2,6- sialyltransferase	Rat liver Galβ1,4GlcNAcα2,6- sialyltransferase
Reducing agent	•	
DTT (1 mM)	1.0	0.9
(1 μM)	1.1	1.2
Mercaptoethanol (1 mM)	1.1	1.1
(1 μM)	1.0	1.1
Detergent		
Triton x-100 (1%)	1.5	0.8
(0.5%)	1.4	1,4
(0.1%)	1.3	1.3
Divalent cations		
MgCl ₂ (5 mM)	· 11 ·	1.0
MnCl ₂ (5 mM)	·13	1.1
EDTA (5 mM)	1.7	0.9

The method of the present invention was specifically explained above referring to the examples relating to the Galβ1,4GlcNAcα2,6-sialyltransferase. However, the method of the present invention is not limited to these examples. As described above, unlike other glycosyltransferases, sialyltransferases share highly conserved regions (sialylmotif, Livingston, B.D. and Paulson, J.C., J. Biol. Chem.,268, 11504-11507, 1993), and all of the sialyltransferases are considered to have similar higher-order structures (Drickamer, K., Glycobiology, 3, 2-3, 1993). Therefore, it is readily understood by those skilled in the art that the renaturation procedure disclosed in the above examples for Galβ1,4GlcNAcα2,6-sialyltransferase can be applied to renaturations of other sialyltransferases to achieve the same advantageous effects. Furthermore, those skilled in the art will be able to choose optimum renaturing conditions, not only for Galβ1,4GlcNAcα2,6-sialyltransferase but for other sialyltransferases, by modifying or altering the processes disclosed in the specification.

The regents and samples used in the above example (D) were as follows. Rat liver Galβ1,4GlcNAcα2,6-sialyttransferase, fetuin, asialo-fetuin, bovine submaxillary mucin, α1-acid glycoprotein, galactose β1,3-N-acetylgalactosamine, lacto N-tetraose and N-acetyllactosamine were obtained from Sigma (St. Louis, USA). Urea was purchased from Wako Pure Chemicals (Osaka, Japan) and a solution was prepared just before use. CMP-[¹⁴C]NeuAc (11 GBq/mmole) was obtained from Amersham (U.K). Bovine submaxillary asialo-mucin and asialo-α1-acid glycoprotein were obtained by mild acid treatment of corresponding glycoproteins. N-acetylgalactosamin e β1,4-galactose was a kind gift from Dr. Kajimoto (The institute of Physical and Chemical Research, RIKEN, Wako-shi, Saitama-ken, Japan). Pyridylamino oligosaccharides (PA-sugar 001, 021, 022 and 023) were obtained from Takara (Kyoto, Japan). Protein concentrations

were determined with a BCA protein assay kit (Pierce) using bovine serum albumin as the standard. Dialysis tubing (20/32) was from Viskase.

Industrial applicability

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The novel GalNAco2,6-sialytransferases P-B1 and P-B3, and proteins which contain a polypeptide part as being the active domain of said enzymes and are released extracellularly provided by the present invention are useful as, for example, reagents for introducing human type sugar-chain to proteins and medicament for treating hereditary diseases lacking human-specific sugar chains. In addition, they can be used as drugs for inhibiting tumor metastases, preventing viral infection, and controlling inflammatory reaction. Furthermore, the method of the present invention is useful when a large quantity of a sialytransferase is expressed in microorganisms, since it enables a mass recovery of the enzyme with highly restored activity from aggregate or precipitate inside the cells.

SEQUENCE LISTING

5	(2) INFORMATION OF SEQ ID NO:1:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 2027	
10	(B) TYPE: nucleic acid	
	(C) STRANDNESS: double	
	(D) TOPOLOGY: linear	
15	(ii) MOLECULE TYPE: cDNA to mRNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
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	MET Gly Phe Leu Ile Arg Arg Leu Pro Lys Asp Ser Arg Ile Phe 15	
30	CGT TGG CTC CTT ATT TTA ACA GTC TTT TCC TTC ATC ATT ACT AGT 90	
	Arg Trp Leu Leu Ile Leu Thr Val Phe Ser Phe Ile Ile Thr Ser 30	
•	·	
35 .	TTT AGC GCC TTG TTT GGC ATG GAG AAA AGC ATT TTC AGG CAG CTC 135	
	Phe Ser Ala Leu Phe Gly MET Glu Lys Ser Ile Phe Arg Gln Leu 45	
0	AAG ATT TAC CAA AGC ATT GCA CAT ATG CTA CAA GTG GAC ACC CAA 180	
40	Lys Ile Tyr Gln Ser Ile Ala His MET Leu Gln Val Asp Thr Gln 60	
45	GAT CAG CAA GGT TCA AAC TAT TCT GCT AAT GGG AGA ATT TCA AAG 225	
	Asp Gln Gln Gly Ser Asn Tyr Ser Ala Asn Gly Arg Ile Ser Lys 75	
50	GTT GGT TTG GAG AGA GAC ATT GCA TGG CTC GAA CTG AAT ACT GCT 270	
•	Val Gly Leu Glu Arg Asp Ile Ala Trp Leu Glu Leu Asn Thr Ala 90	

22

	GIG	AGT	ACA	CCA	AGT	GGG	GAA	GGG	AAG	GAA	GAG	CAG	AAG	AAA	ACA	315
5	Val	Ser	Thr	Pro	Ser	Gly	Ģlu	Gly	Lys	Glu	Glu	Gln	Lys:	Lys	Thr	105
	GTG	AAA	CCA	GTT	GCC	AAG	GTG	GAA	GAA	GCC	AAG	GAG	AAA	GTG	ACT	360
10	Val	Lys	Pro	Val	Ala	Lys	Val	Glu	Glu	Ala	Lys	Glu	Lys	Val	Thr	120
	GTG	AAA	CCA	TTC	CCT	GAG	GTG	ATG	GGG	ATC	ACA	AAT	ACA	ACA	GCA	405
15				Phe												135
	тса	ACA	GCC	тст	GTG	GTG	GAG	AGA	ACA	AAG	GAG	AAA	ACA	ACA	GCG	450
20				Ser												150
		co's	com	001	000	cmc	ccc	CNA	com	C M TO		110	1Ć1	3.03		405
25 .				Pro			•								ACG Thr	495 165
30				CCC												540 180
	•									_		_				
35	AAA	CCA	TCC	TTT	GGG	atg	AAG	GTA	GCT	CAT	GCA-	AAC	AGC	ACA	TCC	585
	Lys	Pro	Ser	Phe	Gly	MET	Lys	Val	Ala	His	Ala	Asn	Ser	Thr	Ser	195
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	Ile	Arg	Pro _.	V al	Thr	Gln	Ala	Ala	Thr	Val	Thr	Glu	Lys	Lys	Lys	225
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	Leu	Arg	Ala	Ala	Asp	Phe	Lys	Thr	Glu	Pro	Gln	Trp	Asp	Phe	Asp	240

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5	Asp	Glu	Tyr	Ile	Leu	Asp	Ser	Ser	Ser	Pro	Val	Ser	Thr	Cys	Ser	· 2	255
	GAA	TCA	GTG	AGA	GCC	AAG	GCT	GCC	AAG	TCT	GAC	TGG	CTG	Cga	GAT	8	10
10	Glu	Ser	Val	Arg	Ala	Lys	Ala	Ala	Lys	Ser	Asp	Trp	Leu	Arg	Asp		70
	CTT	TTC	CTG	CCG	AAC	ATC	ACA	CTC	TTC	ATA	GAC	AAG	agt	TAC	TTC	. 8	55
15	Leu	Phe	Leu	Pro	Asn	Ile	Thr	Leu	Phe	Ile	Asp	Lys	Ser	Tyr	Phe	. 2	85
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25	Gly	Phe	MET	Glu	Leu	Asn	Tyr	Ser	Leu	Val	Glu	Glu	Val	MET	Ser	3	315
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<i>3</i> 5	AGC	AGC	AAC	GTG	TCA	ACG	TGC	ATC	AGC	TGT	- GCT	GTT	GTG	GGG	AAT	. 10	35
. •	Ser	Ser	Asn	Val	Ser	Thr	Cys	Ile	Ser	Суѕ	Ala	Val	Val	Gly	Asn	3	345
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20 .	Lys	Asp	Ile	Arg	Lys	Gly	Phe	Leu	Asn	Tyr	Tyr	Gly	Arg	Arg	Pro	450
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25	Arg	Glu	Arg	Phe	Asp	Glu	Asp	Phe	Thr	MET	Asn	Lys	Tyr	Leu	Val	465
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35						AAG										1485 495
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40						CTG Leu										1530 · 510
	1111	GIY	nia	Dea	ped	nea	Deu		n1a	DCG	1143	Deu	Cla	nap	ura	7 310
45						TAC Tyr										1575 525
		~~*		-1-	1	-1-		_,		1			-1-	-2-		•==
50						AAG Lys										1620 540
			-1-	- 2 - 3		-3-		•	•					•		

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Asp	Glu	Asn	Ile	MET	Lys	Leu	Tyr	Gln	Arg	Ser					566
	GAG	GGCC	ATT	GCCT	YGGGA	AA :	rctc?	ACAC	C A	CCTC	ATGGG	GAA	CAGA	IAGA	1760
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	CCA	CAGA	GAT	ATTT	CCCT	CC :	rttg <i>i</i>	TATO	TT	ratti	TCTC	ACA	ACAC	TTC	1860
	CTA	TAAA	GTG	CATA	TTCT	AC A	AGACO	CAAGO	G A	ACAGI	raggg	AAA	AGTG	CCT	1910
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	CAG	AATG	CTG	GACA	TTGT	AC C	TCTI	GCTG	T GC	GTTC	CCCT	GGC	TGCA	GAC	2410
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(2) INFORMATION OF SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1294

	(B) TYPE: nucleic acid (C) STRANDNESS: double																
•			(c) s	TRAN	DNES	s: d	ouble	e			•					
<i>5</i> .			(1	D) T	OPOL	OGY:	·lin	ear									
		(i	i) M	OLEC	ULE :	TYPE	: cD	NA to	o mRi	NA							
		(x:	i) s	EQUE	NCE I	DESC	RIPT	ION:	SEQ	ID I	NO: 2	:					
10																	
	(-43)		CTG (CAGG	STTT:	et a	MIT.	raat:	r TT	CTTT	CAAA	TAC	TCC!	ACC	-1	
15	PstI																
15				•													
•	ATG	AAA	TTC	AGC	TGG	GTC	ATG	TTC	TTC	CTG	ATG	GCA	GTG	GTT	ACA	45	,
20	MET	Lys	Phe	Ser	Trp	Val	MET	Phe	Phe	Leu	MET	Ala	Val	Val	Thr	15	;
	GGG	GTC	AAT	TCA	GAA	TTC	ACT	GAG	CCA	CAG	TGG	Gat	TTT	GAT	GAT ·	90	
25	Gly	Va1	Asn	Ser	Glu	Phe	Thr	Glu	Pro	Gln	Trp	Asp	Phe	Asp	Asp	30	:
					٠												
	GAG	TAC	ATA	CTG	GAT	AGC	TCA	TCT	CCA	GTA	TCG	ACC	TGC	TCT	GAA	135	
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											,						
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35	Ser	Val	Arg	Ala	Lys	Ala	Ala	Lys	Ser	Asp	Trp	Leu	Arg	Asp	Leu	60	
				AAC					•							225	
	Phe	Leu	Pro	Asn	Ile	Thr	Leu	Phe	Ile	Asp	Lys	Ser	Tyr	Phe	Asn	75	
										· 							
45				TGG												270	
	Val	ser	GIU	Trp	ASP	Arg	Dea	GIU	HIS	Pne	Ald	Pro	Pro	70	GIÅ	90	
				00										,,			
50	Junc	ATY:	GAG	CTG	ልልጥ	ТАС	TCA	CIVE	GT2	GAA	GAA	GTY	ATC	тса	ccc	315	•
															Arg	105	
		5				-1-											
55	-																

	CIG	CCI	CCA	MI	CCC	Cnc	CAG	CAG	CIG	CIC	CIG	GCC	AAC	NOI	AGC	200
5	Leu	Pro	Pro	Asn	Pro	His	Gln	Gln	Leu	Leu	Leu	Ala	Asn	Ser	Ser	120
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10	Ser	Asn	Val	Ser	Thr	Cys	Ile	Ser	Cys	Ala	Val	Val	Gly	Asn _.	Gly	135
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	GAC	TAT	GTG	TTC	CGG	GTG	AGC	GGG _.	GCT	GTA	ATC	AAA	GGT	TAC	GAA	495
20	Asp	Tyr	Val	Phe	Arg	Val	Ser	Gly	Ala	Val	Ile ·	Lys	Gly	Tyr	Glu	165
8	AAG	gat	GTG	GGA	ACA	AAA	ACC	TCC	TTC	TAC	GGA _.	TTC	ACA	GCG	TAC	540
25	Lys	Asp	Val	Gly	Thr	Lys	Thr	Ser	Phe	Tyr	Gly	Phe	Thr	Ala	Tyr	180
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9. 9.5	AAG	ATC	CCA	CAG	GGG.	AAG	CAT	ATC	AGA	TAC	ATT	CAC	TTC	CTG	GAG	630
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40	GCA	GTT	AGA	GAC	TAT	GAG	TGG	CTG	AAG	GCT	CTT	CTG	TTG	GAC	AAG	675
	Ala	Val	Arg	Asp	Tyr	Glu	Trp	Leu	Lys	Ala	Leu	Leu	Leu	Asp	Lys	225
45 [.]	GAT	ATC	AGG	AAA	GGA	TTC	CTG	AAC	TAC	TAT	GGG	CGA	AGG	CCC	CGG	720
·	Asp	Ile	Arg	Lys	Gly	Phe	Leu	Asn	Tyr	Tyr	Gly	Arg	Arg	Pro	Arg	240
50 .	GAG	AGA	TTC	GAT	gaa	gat	TTC	ACA	ATG	AAT	AAG	TAC	CTG	gta	GCT	765
	Glu	Arg	Phe	Asp	Glu	Asp	Phe	Thr	MET	Asn	Lys	Tyr	Leu	Val	Ala	255

	CAC CCT GAT TTC CTC AGA TAC TTG AAA AAC AGG TTC TTA AAA TCT	810
5	His Pro Asp Phe Leu Arg Tyr Leu Lys Asn Arg Phe Leu Lys Ser	270
	AAA AAT CTG CAA AAG CCC TAC TGG CGG CTG TAC AGA CCC ACA ACA	855
o	Lys Asn Leu Gln Lys Pro Tyr Trp Arg Leu Tyr Arg Pro Thr Thr	285
	GGA GCC CTC CTG CTG ACT GCC CTG CAT CTC TGT GAC CGG GTG	900
5	Gly Ala Leu Leu Leu Thr Ala Leu His Leu Cys Asp Arg Val	300
	AGT GCC TAT GGC TAC ATC ACA GAA GGT CAC CAG AAG TAC TCG GAT	945
20	Ser Ala Tyr Gly Tyr Ile Thr Glu Gly His Gln Lys Tyr Ser Asp	315
	CAC TAC TAT GAC AAG GAG TGG AAA CGC CTG GTC TTC TAC GTT AAC	990
95 ·	His Tyr Tyr Asp Lys Glu Trp Lys Arg Leu Val Phe Tyr Val Asn	330
•	CAT GAC TTC AAC TTG GAG AAG CAG GTG TGG AAA AGG CTT CAT GAT	1035
	His Asp Phe Asn Leu Glu Lys Gln Val Trp Lys Arg Leu His Asp	345
15	GAG AAC ATC ATG AAG CTC TAC CAG AGA TCC TGA CAGTGTGCCGAG	1080
•	Glu Asn Ile MET Lys Leu Tyr Gln Arg Ser	360
o	GGCCATTGCC TGGGAAATCT CAACAGCACC TCATGGGGAA CAGAAGAGGA	1175
	CCTCGGAAGC CAGGGTTAGC TCTGGACTTC CAGGCCCAGC TTCAGCTCCA	1225
	CAGAGATATT TCCCTCCTTT GATATC	1251
5	ECORV	

(2) INFORMATION OF SEQ ID NO:3:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1666

(B) TYPE: nucleic acid

	(C) STRANDNESS: double
5	(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: cDNA to mRNA
	(vi) ORIGINAL SOURCE:
10	(A) ORGANISMS: G. gallus (chicken)
	(D) OTHER INFORMATION: CDS 1-1212
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
	-384-TCTTTTTGTCATCAGTGTAA -361
	ATAGGAAGAGCACAAAGTCATTTCTTTCTGCCAATCGCCTTGTGACTCCTTCCGTACATA -301
20	TACACGTGTTGTACAGTATGCTTAAACAGTCCTTGATGAGGTCATCGCTTATTTTTGTTC -241
	TTTTCTGTGTCATAAGAGTTTGGGTTCGCCGCGAATTCGCGGCTAGCCTTGGAGAAGCAG -183
	CGAGTCTGAACCAGTCCGCCCAGCGCCTCCTCCTCCGGCTCACACCCTCCTTCCT
25	GCTCCTCGGAACATCCATCGCTCCGTCCGTCCATCCGTGCGTCCATCCCGGCTGCGGGGA -63
	GCAGCGGAGCGCCGCTTCGGATCCACGCGGACGCGGACCCAACCCGGAGAGCAGCG
30 -	
	ATG GGT TCC CCC CGC TGG AAG CGT TTC TGC TTC TTG CTC CTC GCA 45
	MET Gly Ser Pro Arg Trp Lys Arg Phe Cys Phe Leu Leu Leu Ala 15
35	·
	GCC TTC ACC TCG TCC CTT CTG CTC TAC GGG CAC TAC TAC GCT ACG 90
	Ala Phe Thr Ser Ser Leu Leu Leu Tyr Gly His Tyr Tyr Ala Thr 30
40	
	GTG GAC GTG CGC AGC GGC CCG AGG GTC GTG ACC AGC CTG CTG CAG 135
•	Val Asp Val Arg Ser Gly Pro Arg Val Val Thr Ser Leu Leu Gln 45
45	·
	CCA GAG CTG CTG CTG GTC CGC CCA GAC ACC CCA CAC CCA GAC 180
	Pro Glu Leu Leu Phe Leu Val Arg Pro Asp Thr Pro His Pro Asp 60
50	
	AAC AGC CAC CAC AAG GAG CTC AGA GGG ACT GTG AAG AGC AGG GAG 225

	ASN	ser	HIS	HIS	rys	GIU	reu	Arg	GIŞ	THE	vai	гÃ2	ser	Arg	GIU	/5
	TTC	TTC	TCC	CAA	CCA	TCC	TCA	GAG	CTG	GAG	AAG	ccc	AAA	ccc	AGT	270
	Phe	Phe	Ser	Gln	Pro	Ser	Ser	Glu	Leu	Glu	Lys	Pro	Lys	Pro	Ser	90
o																
	GGA	AAG	CAG	ccc	ACC	CCG	TGC	CCC	CGC	TCG	GTG	GCA	GCC	ACG	GCG .	315
•	Gly	Lys	Gln	Pro	Thr	Pro	Cys	Pro	Arg	Ser	Val	Ala	Ala	Thr	Ala	105
5								-i-					.		222	260
					ACG											360
	rås	AIA	ASP	Pro	Thr	Pne	GIĀ	GIU	rea	Pne	GIN	Pne	иер	116	Pro	120
0	GTG	CTG	ATG	TGG	GĄС	CAA	CAC	TTC	AAC	CCT	GAG	ACG	TGG	GAC	AGG	405
					Asp											135
5				_	_											
	CTG	AAG	GCA	CGA	CGC	GTC	CCA	TAC	GGC	TGG	CAG	GGT	TTG	TCC	CAA	450
	Leu	Lys	Ala	Arg	Arg	Val	Pro	Tyr	Gly	Trp	Gln	Gly	Leu	Ser	Gln	150
o						٠										
	GCA	GCT	GTC	GGC	AGC	ÄCC	CTG	CGT	CTC	CTT	AAC	ACC	TCC	TCC	AAC	495
	Ala	Ala	Val	Gly	Ser	Thr	Leu	Arg	Leu	Leu	Asn	Thr	Ser	Ser	Àsn	165
5	•								,							
					GAC											540
	Thr	Arg	Leu	Phe	Asp	Arg	His	Leu	Phe	Pro	Gly	Gly	Cys	Ile	Arg	180
0	mcm.	ccc	CTVC	CTC.	GGC	3300	ccc	CCN	አጥሮ	ርቁር	እአሮ	ccc	ም ር እ	ccc	CNC	585
					Gly											195
5	0,15		,,,,	,,,	0.1							,	-	,		
	GGC	CGG	GCC	ATC	GAC	GCA	CAT	GAT	TTG	GTC	TTC	AGG	CTG	AAC	GGG	630
	Gly	Arg	Ala	Ile	Asp	Ala	His	Asp	Leu	Val	Phe	Arg	Leu	Asn	Gly	210
ro	GCC	ATC	ACC	AAA	GGC	TTT	GAG	GAG	GAT	GTT	GGG	AGC	AAG	GTT	TCG	675
	Ala	Ile	Thr	Lys	Gly	Phe	Glu	Glu	Asp	Val	Gly	Ser	Lys	Val	Ser	225

	TTC	TAC	GGC	TTC	ACG	GTG	AAC	ACC	ATG	AAG	AAC	TCA	CTC	ATT	GCC	720
5	Phe	Tyr	Gly	Phe	Thr	Val	Asn	Thr	Met	Lys	Asn	Ser	Leu	Ile	Ala	240
	TAT	GAG	GCG	TAT	GGC	TTC	ACC	CGG	ACA	CCG	CAG	GGC	AAG	GAC	CTG	765
10	Tyr	Glu	Ala	Tyr	Gly	Phe	Thr	Arg	Thr	Pro	Gln	Gly	Lys	Asp	Leu	255
•	AAG	TÁC	ATC	TTC	ATC	ccc	TCG	GAC	GCA	CGC	GAC	TAC	ATC	ATG	CTG	. 810
15 .	Lys	Tyr.	Ile	Phe	Ile	Pro	Ser	Asp	Ala	Arg	Asp	Tyr	Ile	Met	Leu	270
20				ATT												855
	Arg	Ser	Ala	Ile	Gln	Gly	Ser	Pro	Val	Pro	Glu	Gly	Leu	Asp	Lys	285
25	GGC	GAC	GAG	CCA	CAG	AAG	TAT	TTT	GGA	CTG	GAG	GCA	TCT	GCG	GAG	900
	Gly	Asp	Glu	Pro	Gln	Lys	Tyr	Phe	Gly	Leu	Glu	Ala	Ser	Ala	Glu	3,0,0
30	AAG	TTC.	AAG	CTG	CTG	CAT	CCC	GAT	TTC	TTG	CAT	TAĊ	CTG	ACA ⁻	ACC	945
	Lys	Phe	Lys	Leu	Leu	His	Pro	Asp	Phe	Leu	His	Tyr	Leu	Thr	Thr	315
35	AGG	TTC	CTG	AGG	TCA	GAG	CTC	CTG	GAC	atg	CAG	TAC	GGC	CAC	CTC	990
:	Arg	Phe	Leu	Arg	Ser	Glu	Leu	Leu	Asp	Met	Gln	Tyr	Gly	His	Leu	330
40	TAC	ATG	ccc	AGC	ACT	GGG	GCA	CTC	ATG	CTG	CTG	ACA	GCA	CTG	CAC	1035
	Tyr	Met	Pro	Ser	Thr	Gly	Ala	Leu	Met	Leu	Leu	Thr	Ala	Leu	His	345
45			.•													
• •				CAG												1080
	Thr	Cys	Asp	Gln	Val	Ser	Ala	Tyr	Gly	Phe	Ile	Thr	Ala	Asn	Tyr	360
50	GAG	CAG	TTC	TCC	GAC	CAT	TAC	TAC	GAG	CCA	GAG	AAG	AAG	CCA	CTG	1125
	Glu	Gln	Phe	Ser	Asp	His	Tyr	Tyr	Glu	Pro	Glu	Lys	Lys	Pro	Leu	375

	GTG TTC TAC GCC AAC CAC GAC ATG CTG CTG GAA GCA GAG CTG TGG	1170
	Val Phe Tyr Ala Asn His Asp Met Leu Leu Glu Ala Glu Leu Trp	390
5		
	AGG AGT TTG CAC CGG GCG GGG ATC ATG GAG CTG TAC CAG CGG TGA	1215
	Arg Ser Leu His Arg Ala Gly Ile Met Glu Leu Tyr Gln Arg	404
10		
	GGGCAGCGCAGTCCCACTGCAAGGACTCTCAATGCAACGCAGAAGCGGTTCTCCTCTTTC	1275
15	CTGAAGGGCTCCTTCTGTCCCTGGAGGGCTCTCCCACACTGGCGGGCCAGCCTGAGGAGC	1335
	AGGGCCTGCAGCTGACAGCCAGAGCAAAGGTGGTGCAGGGCGAGCCAAGGCTGGCAGG	1395
	GAAATACTGCAACTCCTCAGGGCCCTTCAGCATCTTATTTGTGACTCTGAGACTGAGCAC	1455
20	GGCCTTGGGGAGCCTCCGCACGTGGCTGTGAGCTCCTGATGCCATGAGAATGTCTGTGGG	1515
	GTGGCAGCAGCCCCTGGGAAGCACAGTGTTCATGTGCAGGTGGGGCACAGTGGTGCTGGA	1575
	AGGGGATGCTGGAGAAGCATACATCTGACAGACCTCACTTCTTGGAACTTCCTGGAGTTG	1635
25	CAGCCTCGAAGTCACGCTGGGTAGGCTGCAG	1666
	•	
	(2) INFORMATION OF SEQ ID NO:4:	
30	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 1152	
	(B) TYPE: nucleic acid	
35	(C) STRANDNESS: double	
	(D) TOPOLOGY: linear	
0.	(ii) MOLECULE TYPE: DNA	
	(vi) ORIGINAL SOURCE:	
	(A) ORGANISMS: mouse	
45	(D) OTHER INFORMATION: 1-1128 sialyltransferase in so	luble
	form	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
50		
	(-6) TGGCAT	-1
	N	deI

	ATG	GGG	AGC	GAC	TAT	GAG	GCT	CIT	ACA	116	ÇAA	GCC	AAG	GTA	TTC	45
	MET	Gly	Ser	Asp	Tyr	Glu	Ala	Leu	Thr	Leu	Gln	Ala	Lys	Val	Phe	15
	CAG	ATG	CCG	AAG	AGC	CAG	GAG	AAA	GTG	GCC	GTG	GGG	CCT	GCT	ccc	90
0	Gln	MET	Pro	Lys	Ser	Gln	Glu	Lys	Val	Ala	Val	Gly	Pro	Ala	Pro	30
	CAG	CCT	GTG.	•	ጥሮኔ	AAC	ACC	222	CAA	GAC	ርር ጥ	AAG	CAA	ccc	Custr	135
··5						Asn										45
	CAG	ATC	CTC	AGT	TAC	ccc	AGG	GTC	ACA	GCC	AAG	GTC	AAG	CCA	CAG	180
20	Gln	Ile	Leu	Ser	Tyr	Pro	Arg	Val	Thr	Ala	Lys	Val	Lys	Pro	Gln	60
	ccc	TCC	TTG	CAG	GTG	TGG	GAC	AAG	GAC	TCC	ACA	TAC	TCA	AAA	CTT	225
25						Trp	•									75
•																
30	AAC	CCC	AGG	CTG	CTG	AAG	ATC	TGG	AGG	AAC	TAT ·	CTG	AAC	atg	AAT	270
	Asn	Pro	Arg	Leu	Leu	Lys	Ile	Trp	Arg	Asn	Tyr	Leu	Asn	MET	Asn	90
	AAA	TAT	AAA	GTG	TCC	TAC	 AAG	GGG	CCG	GGA	CCA	GGA	GTC	AGG	TTC	315
35	Lys	Tyr	Lys	Val	Ser	Tyr	Lys	Gly	Pro	Gly	Pro	Gly	Val	Arg	Phe	105
							ż				•					
10						CGC	_		_		_			_		360
	Ser	Val	Glu	Gly	Leu	Arg	Cys	His	Leu	Arg	Asp	His	Val	Asn	Val	120
15	TCT	ATG	ATA	GAG	GCC	ACA	GAT	TCT	ccc	TTC	AAC	ACC	ACT	GAA	TGG	405
	Ser	MET	Ile	Glu	Ala	Thr	Asp	Ser	Pro	Phe	Asn	Thr	Thr	Glu	Trp	135
									٠							
50						AAA Lys										. 450 150
	OIU	GIÅ		Den	FEO	rås	GIU	THE	FILE	wrd	HIL	nys	WIG	GIÅ	FIO	720

•	TGC	ACA	AAG	TGT	GCC	GTC	GTG	TCT	TCT	GCA	GGA	TCT	CTG	AAG	AAC	495
5	Cys	Thr	Lys	Cys	Ala	Val	Val	Ser	Ser	Ala	Gly	Ser	Leu	Lys	Asn	165
	TCC	CAG	CTG	GGT	CGA	GAG	ATT	gat	AAT	CAT	GAT	GCG	GTC	CTG	AGG	540
10	Ser	Gln	Leu	Gly	Arg	Glu	Ile	Asp	Asn	His	Asp	Ala	Val	Leu	Arg	180
												٠				
	TTT	AAT	GGG	GCA	CCT	ACA	GAC	AAC	TTC	CAA	CAG	GAT	GTG	GGC	ACA	585
15	Phe	Asn	Gly	Ala	Pro	Thr	Asp	Asn	Phe	Gln	Gln	Asp	Val	Gly	Thr	195
20	AAA	ACT	ACC	ATC	CGC	CTA	GTG	AAC	TCT	CAG	TTA	GTC	ACC	ACA	GAA	630
	Lys	Thr	Thr	Ile	Arg	Leu	Val	Asn	Ser	Gln	Leu	Val	Thr	Thr	Glu	210
•				٠.												
25	AAG	CGC	TTC	CTG	AAG	GAC	agt	TTG	TAC	ACC	GAA	GGA	ATC	CTG	ATT	675
	Lys	Arg	Phe	Leu	Lys	Asp	Ser	Leu	Tyr	Thr	Glu	Gly	Ile	Leu	Ile	225
						٠.										
30	CTG	TGG	GAC	CCA	TCT	GTG	TAT	CAT	GCA	GAC	ATT	CCG	CAG	TGG	TAT	720
	Leu	Trp	Asp	Pro	Ser	Val	Tyr	His	Ala	Asp	Ile	Pro	Gln	Trp	Tyr	240
	CAG	AAG	CCA	GAC	TAC	AAC	TTC	TTC	GAA	ACC	TAT	AAG	agt	TAC	CGA	765
35	Gln	Lys	Pro	Asp	Tyr	Asn	Phe	Phe	Glu	Thr	Tyr	Lys	Ser	Tyr	Arg	255
40	•			CCC												810
	Arg	Leu	His	Pro	Ser	Gln	Pro	Phe	Tyr	Ile	Leu	Lys	Pro	Gln	MET	270
45				CTA						•						855
	Pro	Trp	Glu	Leu	Trp	Asp	Ile	Ile	Gln	Glu	Ile	Ser	Pro	Asp	Leu	285
50				AAT												900
	Ile	Gln	Pro	Asn	Pro	Pro	Ser	Şer	Gly	MET	Leu	Gly	Ile	Ile	Ile	300

	ATG	ATG	ACG	CTG	TGT	GAC	CAA	GTT	GAT	ATT	TAC	GAG	TTC	CTC	CĊA	945
5	MET	MET	Thr	Leu	Cys	Asp	Gln	Val	Asp	Ile	Tyr	Glu	Phe	Leu	Pro	315
	тсс	AAG	CGC	AAG	ACA	GAT	GTG	TGC	TAC	TAT	CAC	CAG	AAG	TTC	TTT	990
10	Ser	Lys	Arg	Lys	Thr	Asp	Val	Cys	Tyr	Tyr	His	Gln ·	Lys	Phe	Phe	330
			200	maa	100	a m/C	CCM	ccc	ma C	Cam	ccc	ריייר		ጥ ሞር	CAG	1035
15	GAC	AGC	GCC	TGC	ACG	AIG	GGT	GCC	TAC	CAI	CCG	CIC	CIC	ŢTC	GAG	1033
	Asp	Ser	Ala	Cys	Thr	MET	Gly	Ala	Tyr	His	Pro	Leu	Leu	Phe	Glu	345
20	AAG	AAT	ATG	GTG	AAG	CAT	CTC	AAT	GAG	GGA	ACA	GAT	GAA	GAC	ATT	1080
	Ĺys	Asn	MET	Val	Lys'	His	Leu	Asn	Glu	Gly	Thr	Asp	Glu	Asp	Ile	360
<i>25</i>												*				
20	TAT	TTG	TTT	GGG	AAA	GCT	ACC	CTG	TCT	GGC	TTC	CGG	AAC	AAT	CGC.	1125
	Tyr	Leu	Phe	Gly	Lys	Ala	Thr	Leu	Ser	Gly	Phe	Arg	Asn	Asn	Arg	375
30																
	TGT	TGA	GCC	AGGG	ATCC:	CAT										1146
	Cys			Ва	amHI									•	•	376
35	-										•					

40 Claims

- 1. GalNAca2,6-sialytransferase P-B1 characterized by the amino acid sequence defined by SEQ ID No.1 of the sequence listings.
- A GalNAco2,6-sialytransferase P-B1 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B1.
 - 3. The GalNAcα2,6-sialyltransferase gene according to claim 2 characterized by the nucleotide sequence of from nucleotides 1 to 1698 defined by SEQ ID No.1 of the sequence listings.
 - GalNAcα2,6-sialytransferase P-B3 characterized by the amino acid sequence defined by SEQ ID No.3 of the sequence listings.
- 5. A GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the amino acid sequence of GalNAco2,6-sialytransferase P-B3 gene encoding the acid sequence of GalNAco2,6-sialytransferase of GalNAco2,6-sialytr
 - 6. The GalNAco2,6-sialyltransferase P-B3 gene according to claim 5 characterized by the nucleotide sequence of from nucleotides 1 to 1212 defined by SEQ ID No.3 of the sequence listings.

- A recombinant vector comprising a GalNAcα2,6-sialyltransferase gene encoding the amino acid sequence of the GalNAcα2,6-sialyltransferase P-B1.
- The recombinant vector according to daim 7 which is plasmid ACEB-3034.

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- A recombinant vector comprising a GalNAca2,6-sialyltransferase gene encoding the amino acid sequence of the GalNAca2,6-sialyltransferase P-B3.
- 10. The recombinant vector of claim 9 which is plasmid ACEB3-T20 or plasmid pcDB3ST.
- A transformant which is being transformed with a recombinant vector comprising a GalNAcα2,6-sialyltransferase gene encoding the amino acid sequence of the GalNAcα2,6-sialyltransferase P-B1.
- 12. A transformant which is being transformed with a recombinant vector comprising a GalNAco2,6-sialyltransferase gene encoding the amino acid sequence of the GalNAco2,6-sialyltransferase P-B3.
 - 13. An extracellularly releasable protein catalyzing GalNAco2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAco2,6-sialytransferase P-B1 or P-B3 together with a signal peptide.
- 20 14. The protein according to claim 13, wherein the active domain of the GalNAcα2,6-sialyltransferase P-B1 is the polypeptide characterized by amino acids of from 233 to 566 defined by SEQ ID No.1 of the sequence listings.
 - 15. The protein according to daim 13 which is the protein SB-690 characterized by the amino acid sequence defined by SEQ ID No.2 of the sequence listings.
 - 16. A gene encoding an extracellularly releasable protein catalyzing GalNAcα2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAcα2,6-sialyltransferase P-B1 or P-B3 together with a signal peptide.
- 30 17. The gene according to claim 16 characterized by the nucleotide sequence of from nucleotides 1 to 1065 defined by SEQ ID No.2 of the sequence listings.
 - 18. A recombinant vector comprising a gene encoding an extracellularly releasable protein catalyzing GalNAcα2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAcα2,6-sialyltransferase P-B1 or P-B3 together with a signal peptide.
 - 19. The recombinant vector according to claim 18 which is plasmid pcDSB-690.
- 20. A transformant which is being transformed with a recombinant vector comprising a gene encoding an extracellularly releasable protein catalyzing GalNAco2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAco2,6-sialytransferase P-B1 or P-B3 together with a signal peptide.
 - 21. A process for preparing an extracellularly releasable protein catalyzing GalNAco2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAco2,6-sialyltransferase P-B1 or P-B3 together with a signal peptide, which comprises the steps of culturing a transformant which is being transformed with a recombinant vector comprising a gene encoding an extracellularly releasable protein catalyzing GalNAco2,6-sialic acid transfer which comprises a polypeptide portion as being an active domain of the GalNAco2,6-sialyltransferase P-B1 or P-B3 together with a signal peptide, and recovering said protein from the culture.
- 50 22. GalNAcα2,6-sialyltransferase.
 - 23. GalNAca2,6-sialyltransferase derived from mammals.
 - 24. A process for preparing a sialyttransferase which comprises the steps of:
 - (a) expressing a sialyltransferase in a microorganism;
 - (b) extracting the sialytransferase using 5 to 9 M urea from proteinic aggregate or precipitate that contains the enzyme and is accumulated inside the microorganism;

- (c) diluting the extract obtained by the step (b) with a renaturation composition to obtain a primary dilution containing 1 to 4 M urea;
- (d) diluting the primary dilution obtained by the step (c) with a renaturation composition to obtain a secondary dilution containing 0.5 to 2 M urea; and
- (e) removing urea from the secondary dilution obtained by the step (d) by dialysis to afford a renatured sialyl-transferase.
- 25. The process according to claim 24, wherein the renaturation composition used in the step (c) comprises 1 to 2 M urea, 20 mM MOPS-NaOH, 0.5M NaCl, 20 mM lactose, and 0.5 mM EDTA (pH 7.0); and the renaturation composition used in the step (d) comprises 20 mM MOPS-NaOH, 0.5 M NaCl, 20 mM lactose, and 0.5 mM EDTA (pH 7.0).
- 26. The process according to claim 24, wherein the sialyltransferase is GalNAcα2,6-sialyltransferase or Galβ1,4GlcNAcα2,6-sialyltransferase.
- 27. A process for preparing a sialytransferase which comprises the steps of:

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- (a) expressing a sialyttransferase in a microorganism;
- (b) extracting the sialytransferase using 8 M urea from proteinic aggregate or precipitate that contains the enzyme and is accumulated inside the microorganism;
- (c) diluting the extract obtained by the step (b) with a renaturation composition, followed by standing the dilution for at least 12 hours at 4 °C to obtain a primary dilution containing 2 to 3 M urea;
- (d) diluting the primary dilution obtained by the step (c) with a renaturation composition, followed by standing the dilution for at least 48 hours to obtain a secondary dilution containing 1 to 2 M urea; and
- (e) removing urea from the secondary dilution obtained by the step (d) by dialysis in the presence of one or more divalent cations to afford a renatured sialyttransferase.
- 28. The process according to claim 27, wherein the renaturation composition used in the step (c) comprises 1 to 2 M urea, 20 mM MOPS-NaOH, 0.5 M NaCl, 20 mM lactose, and 0.5 mM EDTA (pH 7.0); and the renaturation composition used in the step (d) comprises 20 mM MOPS-NaOH, 0.5M NaCl, 20 mM lactose, and 0.5 mM EDTA (pH 7.0).
- 29. A process according to claim 27, wherein the sialyltransferase is GalNAc α 2,6-sialyltransferase or Gal β 1,4GlcNAc α 2,6-sialyltransferase.

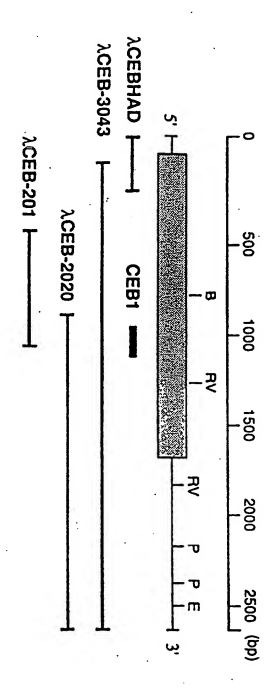
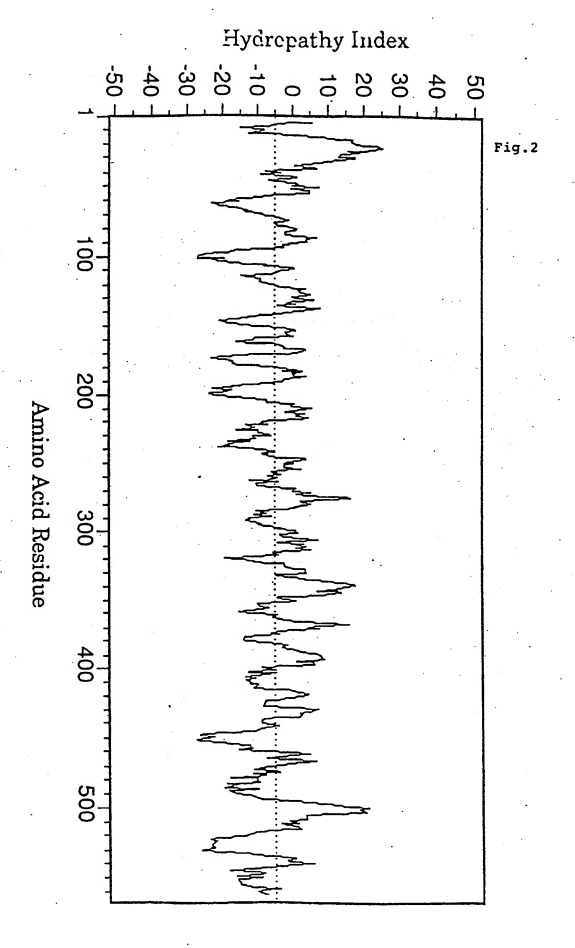


Fig.1



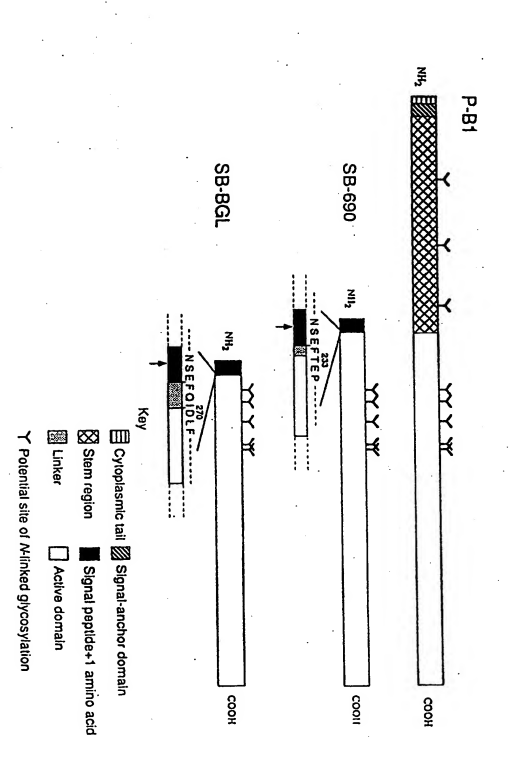


Fig. 4	- 91	-123	-123	-223	-321	
	- MGSPRWKR <u>ecelllaafisslilyg</u> hyyatvdvrsgervvtsllopellflvrddhydhhkelrgtvksreffsopsselekpkpsg 	- KQPTPCPRSVAATAKADPTFGELFQFDIPVLM	- AEEPPASVKAIRPVTQAATVTEKKKLRAADFKTEPQ#DFDDEYILDSSSPVSTCSESVRAKAAKSDWLRDLFLPNITLFIDKSYFNVSEWDRLEHFAPPY	B3 - WDQHFNPETWDRLKARRVPYGWQGLSQAAVGSTLRLFDRHLFPGGCIRCAVVGNGGILNGSRQGRAIDAHDLVFRLNGAITKGFEEDVGSKVSFYGFTVN -223 	– TMKNSLIAYEAYGFTRTPOGKDLKYIFIPSDARDYIMLRSAIQGSPVPEG-LDKGDEPQKYFGLEASAEKFKLL-HPDFLHYLTTRFLRSELLDMQYGHL 	- B3 - YMPSTGALMALTALHTCDQVSAXGFITANYEQFSDHYYEPEKKPLVFYANHDMLLEAELWRSLHRAGIMELYQR -404
	P – B3 P – B1	P – B3 P – B1	P – B3	P – B3 P – B1	P – B3 P – B1	P – B3 P – B1

INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP94/02182

A. CLASSIFICATION OF SUBJECT MATTER Int. Cl ⁶ Cl2N9/10, Cl2N15/54	-										
According to International Patent Gausification (IPC) or to both national dassification and IPC B. FIELDS SEARCHED											
Minimum documentation searched (classification system followed by chanification symbols)											
Int. Cl ⁶ Cl2N9/10, Cl2N15/54											
Documentation scarched other than minimum documentation to the extent that such documents are included in the fields searched											
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)											
CAS ONLINE, BIOSIS, WPI/WPIL											
C. DOCUMENTS CONSIDERED TO BE RELEVANT											
Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim ?	lo.										
PX J. Biol. Chem. Vol. 269, No. 29 (1994), 1-29											
Kurosawa Nobuyuki et al. "Cloning and expression of Gal. beta 1, 3 GalNAc-Specific											
GalNAc alpha 2, 6-sialyltransferase* p. 19048 - 19053	-										
PX J. Biol. Chem. Vol. 269, No. 2 (1994), 1-29											
Kurosawa Nobuyuki et al. "Molecular cloniny and											
expression of GalNAc. alpha 2, 6-sialyltransferase p. 1402-1409	İ										
	ł										
A Annu Rev. Biochem. Vol. 50 (1981), 1-29 p. 733-764											
Further documents are listed in the continuation of Box C. See patent family annex.											
* Special categories of clied documents: "A" document defining the general state of the art which is not considered to be of perfectly relievance to be of perfectly relievance	crity										
"B" earlier document but published on or after the international filling date "A" document of purificalty relevance; the claimed investion trans-											
cited to establish the politication date of another challen are other											
"O" document referring to an oral discioners, we, exhibition or other means or select to ignore an ignorative step when the document means or select the document of the select the select the document of the select the se	2 je										
"?" document published prior to the international (Hing date but later than the priority data claimed "A" document member of the same parent family											
Date of the actual completion of the international search Date of malling of the international search report											
January 25, 1995 (25. 01. 95) February 14, 1995 (14. 02. 95)	١										
Name and mailing address of the ISA/ Authorized officer											
Japanese Patent Office											
Form PCT/ISA/210 (second sheet) (July 1992)											